

STATE OF RHODE ISLAND DEPARTMENT OF HEALTH - OFFICE OF FOOD PROTECTION,
 RM 203, CANNON BLDG, 3 CAPITOL HILL, PROVIDENCE, RI 02908

SAMPLE COLLECTION FORM

LAB # _____

DATE OF COLLECTION 9/14/98		DATE OF ANALYSIS 8/13/98		DATE OF REPORT 8/17/98		LAB SUP PCE	
CONDITION HOT _____ FROZEN _____ COLD _____ OTHER _____		ITEM OYSTERS		PRODUCT CODE/COM		MOP	
SHIP DATE		BRAND NAME		TEMP ESTAB		TEMP REC °C	
DATE OF SHIP		FROM LOT OF		COLLECTED AT Billington Cove Marina		REASON 01	
SHIPPER/PACKER & ADDRESS		ADDRESS		FOLLOW UP SAMPLE #1		SOURCE 10	
COMPLAINT		ORIGINAL CONTAINER YES _____ NO _____		ANALYSIS 13		DATE PURCHASED	
NAME & LOCATION OF STORE WHERE PURCHASED		PRODUCT USED YES _____ NO _____		AMOUNT REMAINING			
HOW STORED FROZEN _____ AMBIENT _____ COLD _____		IMPORT PRODUCT YES _____ NO _____		INTERVIEWED BY		DATE	
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:		TIME		HOW STORED			
SIGNED FROM John Muller		SIGNED TO Matthew Silva		DATE 8/14/98		TIME 1:40	
AGENCY PRINT NAME 164 John Muller		PRINT NAME Matthew Silva					
SIGNED FROM		SIGNED TO					
PRINT NAME		PRINT NAME					
SIGNED FROM		SIGNED TO					
PRINT NAME		PRINT NAME					
1. CEREAL	2. MEAT PROT	3. TOT H ₂ O	4. SOY FL	5. MFOM	6. ADD H ₂ O	7. TOT PROT	8. TOT FAT
%	%	%	%	%	%	%	%
9. SORBATES	10. TBA	11. PFA	12. WT/VOL	13. NaNO ₂	14. EL TROPHO	15. TOTAL SOLIDS	16. HISTAMINE
ppm	ug/mg/100g	ppm	%	ppm	ppm	mg/100g	mg/100g
17. NaCl	18. NH ₄	19. SULFITE	20. Hg	21. TVB	22. DOMOIC ACID	23. TMA	24. VITAMIN A
%	mg/100g	ppm	ppm	mg N/100g	ug/ml	mg/100g	IU/Oz
25. pH	26. BRIX	27. INSECT	28. RODENT	29. BRENZT'S	30. ASCORB'S	31. INDOLE	32. VITAMIN D
	%	%	%	ppm	ppm	ug/100g	IU/Oz
33. FEG INT	34. AFLATOXIN	35. Pb	36. Cd	37. Cu	38. Cr	39. Zn	40. A _v
	ppm	ppm	ppm	ppm	ppm	ppm	
41. Na	42. TSP	43. Ca	44. P	45. Mg	46. GLUTAMIC MSG	47. ACETIC ACID	48. GLUCOSE
mg/100g	mg/100g	mg/100g	mg/100g	mg/100g	%	mg/100g	%
49. FRUCTOSE	50. SUCROSE	51. LACTOSE	52. MALTOSE	53. YEAST	54. MOLD	55. PSP	56. OTHER
%	%	%	%	%	%	%	%
57. PECALAMPN	58. TOT COLIFORMAMPN	59. CL ROT	60. CL PERV	61. CDAG STAPH	62. NON-CDAG STAPH	63. LISTENA	64. CAMPYLO
65. SPC	66. SALMONELLA	67. S CEREUS	68. YERSINA	ORGANOLEPTIC			
COLLECTOR'S NOTES: Morassom Oyster nursery stock. appt. 36nn				LABORATORY NOTES: Fe: 133 ppm			
EVALUATION BY LAB: 1. NOT AN FP VIOLATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)				ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE			
FOOD QUALITY (SAFETY SHELF LIFE & CONSUMER ACCEPTANCE) INVOLVES THE ENTIRE CHAIN OF PRODUCTION PROCESSING & DISTRIBUTION OF FOOD. * SAMPLE WILL BE MAINTAINED FOR THIRTY (30) DAYS AFTER ANALYSIS AND THEN DISPOSED OF BY THE LABORATORY.				CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY			
WHITE - DATA ENTRY		YELLOW - LAB		PINK - FOOD PROTECTION		GOLDENROD - CONSUMER	

Data Collected by Dr. Dale Leavitt, Roger William University

Hard Clam Seed from Warwick Cove Marina

Warwick Cove Upweller																		
Quahog Seed		30-Oct-08		Note: µg/kg = ppb														
Group (n=15)	avg Length (mm)	stdev	avg Live Weight (g)	stdev	avg Soft Tissue Wet Weight (g)	stdev	avg Hg/Soft Tissue Wet Weight (µg/kg)	stdev	avg Cr/Soft Tissue Wet Weight (µg/kg)	stdev	avg Fe/Soft Tissue Wet Weight* (µg/kg)	stdev	avg Ni/Soft Tissue Wet Weight (µg/kg)	stdev	avg Cu/Soft Tissue Wet Weight (µg/kg)	stdev		
1	15.1	2.9	0.967	0.730	0.267	0.238	7.81		0.20		35.57		0.20		8.18			
2	12.6	1.6	0.545	0.202	0.139	0.056	9.41		0.28		34.00		0.22		11.80			
3	13.9	1.2	0.685	0.201	0.182	0.058	8.24		0.26		33.33		0.20		9.30			
Total	13.9	2.2	0.732	0.476	0.196	0.152	8.49	0.83	0.25	0.04	34.30	1.15	0.21	0.01	9.76	1.85		
Group (n=15)					avg Soft Tissue Dry Weight (g)	stdev	avg % Dry Weight	stdev	avg Hg/Soft Tissue Dry Weight (µg/kg)	stdev	avg Cr/Soft Tissue Dry Weight (µg/kg)	stdev	avg Fe/Soft Tissue Dry Weight* (µg/kg)	stdev	avg Ni/Soft Tissue Dry Weight (µg/kg)	stdev	avg Cu/Soft Tissue Dry Weight (µg/kg)	stdev
1					0.041	0.041	14.8%	2.4%	52.75		1.38		240.70		1.37		55.33	
2					0.022	0.008	16.1%	1.5%	58.45		1.73		210.60		1.35		73.09	
3					0.027	0.008	15.2%	1.7%	54.22		1.70		219.20		1.31		81.16	
Total					0.030	0.025	15.4%	2.0%	55.14	2.96	1.60	0.19	223.50	15.50	1.34	0.03	63.19	9.05
Group (n=15)	avg Zn/Soft Tissue Wet Weight* (µg/kg)	stdev	avg As/Soft Tissue Wet Weight (µg/kg)	stdev	avg Se/Soft Tissue Wet Weight (µg/kg)	stdev	avg Sr/Soft Tissue Wet Weight (µg/kg)	stdev	avg Ag/Soft Tissue Wet Weight (µg/kg)	stdev	avg Cd/Soft Tissue Wet Weight (µg/kg)	stdev	avg Pb/Soft Tissue Wet Weight (µg/kg)	stdev				
1	55.10		1.77		0.87		15.28		0.03		0.06		0.31					
2	66.07		1.89		1.29		14.07		0.03		0.05		0.28					
3	55.07		1.55		0.53		11.94		0.03		0.11		0.25					
Total	58.75	6.34	1.77	0.22	0.89	0.38	13.76	1.69	0.03	0.00	0.07	0.03	0.28	0.03				
Group (n=15)	avg Zn/Soft Tissue Dry Weight* (µg/kg)	stdev	avg As/Soft Tissue Dry Weight (µg/kg)	stdev	avg Se/Soft Tissue Dry Weight (µg/kg)	stdev	avg Sr/Soft Tissue Dry Weight (µg/kg)	stdev	avg Ag/Soft Tissue Dry Weight (µg/kg)	stdev	avg Cd/Soft Tissue Dry Weight (µg/kg)	stdev	avg Pb/Soft Tissue Dry Weight (µg/kg)	stdev				
1	372.90		12.01		5.89		103.40		0.22		0.43		2.09					
2	409.30		12.32		7.97		87.14		0.16		0.29		1.76					
3	362.20		10.17		3.47		78.55		0.22		0.69		1.68					
Total	381.47	24.69	11.50	1.16	5.78	2.26	89.70	12.62	0.20	0.04	0.47	0.20	1.84	0.22				



OKATEST

ZE/OA48C

ZE/OA96C

Test for detection of Okadaic Acid-toxins group

**Test para la detección de las toxinas del grupo
del Ácido Okadaico**

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SCOPE

This protocol specifies a method for the quantitative determination of Okadaic Acid (OA) and other carboxylic toxins of the OA group including DTX1, DTX2 and DTX3 by a colorimetric phosphatase inhibition assay. This method is applicable to shellfish species such as mussels, clams, cockle, scallops, etc.

PRINCIPLE

Test based on the phosphatase activity inhibition by OA-toxins group, responsible for diarrhetic shellfish poisoning (DSP).

Phosphatase enzyme PP2A is able to hydrolyse a specific substrate, yielding a product that can be detected at 405 nm. Samples containing toxins from the okadaic acid group will inhibit the enzyme activity proportionally to the amount of toxin contained in the sample. The concentration of toxin in the sample can be calculated using a standard curve.

KIT CONTENTS

	48 Tests Kit	96 Tests Kit
Microtiter plate strips (8 wells per strip)	6	12
Vials of Phosphatase (<i>Phosphatase</i>)	2	4
Set of Okadaic Acid Standards (<i>Okadaic acid 0.5, 0.8, 1.2, 1.8 and 2.8 nM</i>)	1	1
Chromogenic Substrate (<i>Chromogenic Substrate</i>)	1	1
Phosphatase Dilution Buffer (<i>Phosphatase Dilution Buffer</i>)	1	1
Stock Buffer Solution (<i>Stock Buffer Solution</i>)	1	1
Stop Solution (<i>Stop Solution</i>)	1	1
Adhesive film	1	2
Kit instructions	1	1

ADDITIONAL MATERIAL AND REAGENTS NEEDED

- Micropipettes
- Blender (Ultraturax) or mortar and pestle
- Heater at 30°C ± 2 °C (i.e. FX Incubator, Ref ZE/FX, from ZEULAB)
- Microplate reader (wavelength at 405 nm)
- Water bath for 76 ± 2 °C
- Methanol (analytical grade)
- NaOH 2.5 N made by titration, (NaOH of analytical grade)
- HCl 2.5 N made by titration, (HCl of analytical grade)
- Deionised water (grade 2, ISO3696)
- Graded 50 mL centrifuge tubes with screw caps
- Tube shaker
- Centrifuge

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SOLUTIONS

- 1.- **Okadaic Acid Standards:** to make sure these solutions are homogeneous, it is very important to mix well using a vortex, before applying to the plate.
- 2.- **Chromogenic Substrate solution:** The solution contains stabilization resin. Make sure this resin is not added to the microwells. To assure that, it is recommended to transfer the volume needed into a transparent labware (i.e.: test tube or eppendorf) and take the solution from that container to add into the wells. *Note:* Do not use this solution if the absorbance of 90 μL of this solution at 405 nm is over 0.6.
- 3.- **Phosphatase solution:** Add 2.0 mL of phosphatase dilution buffer (*Phosphatase Dilution Buffer*) to one of the phosphatase vials (*Phosphatase*) and dissolve by mixing gently for 1 hour \pm 5 minutes at room temperature (22 ± 2 °C) to ensure that the enzyme is fully hydrated. Do not use the tube shaker at any moment. This solution must be stored under refrigeration if not in use immediately after preparation. Do not use the phosphatase solution for following days. Each enzyme vial contains enough volume for 24 wells. If more than one vial is used in the assay, dissolve each vial as described above, make a pool with the content of the vials and mix gently, by inversion, before use.
*Attention: this reagent is blue and becomes brownish when dissolved. If brownish colour is noticed before hydration, discard this reagent as it could be damaged.
- 4.- **Buffer solution x1:** dilute the *Stock Buffer Solution* included in the kit by mixing 1 volume with 9 volumes of deionised water. Use buffer solution x1 only freshly made, and store under refrigeration if not in use immediately.
- 5.- **2.5 N NaOH:** weigh 100 g of NaOH and add 500 mL of water and dissolve. Transfer to a volumetric flask and add deionised water up to a final volume of 1000 mL.
- 6.- **2.5 N HCl:** add 205 mL of HCl (37 %) to 400 mL of deionised water already contained in a volumetric flask. Make the volume up to 1000 mL with deionised water.

SAMPLES EXTRACTION

The method described below includes a hydrolysis step to detect all toxins forms of okadaic acid (okadaic acid and dinophisistoxins).

- 1.- Clean the shell thoroughly using water
- 2.- Open the shellfish by cutting the adductor muscles.
- 3.- Wash inside the shell thoroughly to remove any dirt.
- 4.- Remove the tissue inside the shell by cutting all the muscles attached to the shell.
- 5.- Place the shellfish tissue in a filter paper for few minutes to remove water in excess.

It is recommended to use graded 50 mL centrifuge tubes with screw caps during the following steps of hydrolysis in order to prevent loses due to labware changes.

- 6.- Mash the shellfish tissue to obtain a representative sample and weigh 5 g. Add 25 mL of Methanol and homogenise the mixture for 2 minutes using a tube shaker.
- 7.- Centrifuge at 2000 g for 10 min at 4 °C. The supernatant (*methanolic extract*) is poured into a centrifuge tube.
- 8.- Take 640 μL of *methanolic extract* and pour into another centrifuge tube.
- 9.- Add 100 μL of 2.5 N NaOH.
- 10.- Seal and heat at 76 ± 2 °C for 40 minutes.

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- 11.- Add 80 μL of 2.5 N HCl (the sample does not need to be cooled down previously).
- 12.- Add up to 20 mL of Buffer solution x1.

TEST PROCEDURE

Warning:

The volume of some reagents used in this assay is small and special attention must be paid when added to the wells:

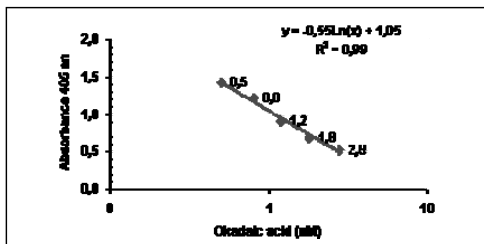
- Make sure the pipettes are calibrated before running the assay.
- Use pipettes according to the volumes to be dispensed. Use pipettes with a maximum pipette volume of 100 or 200 μL .
- Be sure that the incubator's temperature is stabilized before use.

It is recommended to run samples and standards in duplicate.

- 1.- Add 50 μL of samples or standards.
- 2.- Add 70 μL of the Phosphatase Solution to each well. Mix well by gentle tapping on the side of the plate.
- 3.- Cover the plate with the adhesive film provided and incubate for 20 ± 0.5 minutes at 30 ± 2 °C.
- 4.- Remove the adhesive film and add 90 μL of Chromogenic Substrate to each well. Mix well by gently tapping on the side of the plate.
- 5.- Cover the plate with the adhesive film and incubate 30 ± 0.5 minutes at 30 ± 2 °C.
- 6.- Remove the adhesive film and add 70 μL of Stop Solution to each well.
- 7.- Read absorbance of samples and standards at 405 nm.

GRAPHIC REPRESENTATION AND CALCULATIONS OF RESULTS

- 1.- Obtain a standard curve by plotting the absorbance values in a linear y axis and the concentration of okadaic acid in a logarithmic x axis and use a logarithmic fitting as shown in the graphic next page. R^2 has to be greater than or equal to 0.96.



2.- The OA concentration contained in the sample (Cs) is calculated by interpolation into the calibration curve or using the following equation:

$$x = \text{EXP} (y - b/a)$$

Where x is the OA concentration in the sample (Cs) and y the absorbance of the sample.

Note: An Excel worksheet to calculate results is available upon request.

3.- Calculate the diarrhetic shellfish toxins concentration in tissue (Ct) as follows:

$$Ct (\mu\text{g/kg}) = \frac{(Cs (\text{nM}) \times FD \times MW (\text{g/mol}) \times Ve (\text{L}))}{Mt (\text{g})}$$

Ct: toxins concentration in tissue, expressed as equivalents of OA; Cs: toxins concentration in sample; FD: Methanolic extract dilution factor (i.e. 640 $\mu\text{L}/20 \text{ mL} \rightarrow \times 31.25$); MW: Okadaic acid molecular weight = 805; Ve: Methanolic extract volume (0.025L); Mt: Tissue weight (5g).

Example: for OA concentration of 1.5 nM: 1.5 nM \times 31.25 \times 805 g/mol \times 0.025L / 5g = 189 μg OA eq/kg.

NOTE: For samples with OA concentration falling outside the working range (< 0.5 nM or > 2.8 nM), results will be reported as < 0.5 nM (or < 63 $\mu\text{g}/\text{Kg}$) or > 2.8 nM (or > 352 $\mu\text{g}/\text{kg}$), respectively. When sample absorbance is below the value obtained for 2.8 nM the methanolic extract could be diluted up to 1:4 and samples re-tested

STABILITY AND STORAGE

The kit contents must be stored at 4 - 12 °C and protected from light. This kit has a shelf life of 8 months when stored under optimal conditions. See the expiry date on the kit package.

SAFETY

Safety clothing should be worn and skin contact with the reagents avoided. Do not ingest. A SAFETY DATA SHEET is available from your local distributor on request.

***Warning:** Okadaic Acid is toxic. Gloves, mask and other protective clothing must be worn when handling okadaic acid solutions.

REFERENCES

1. Takai, A.; Bialojan, C.; Troschka, M.; Rüegg, J.C. *Smooth muscle myosin phosphatase inhibition and force enhancement by black sponge toxin*. *FEBS Lett.* 1987, 21781-21784.
2. Smienk H., Calvo D., Razquin P., Domínguez E. & Mata L. *Single Laboratory Validation of A Ready-to-Use Phosphatase Inhibition Assay for Detection of Okadaic Acid Toxins*. *Toxins*, 2012, 5, 339-352.
3. Smienk H., Domínguez E., Rodríguez-Velasco M.L. Clarke D., Katrin K., Katikou P., Cabado A.G., Otero A., Vieties J.M. Razquin P., and Mata L. *Quantitative Determination of the Okadaic Acid Toxins Group by a Colorimetric Phosphatase Inhibition Assay: Interlaboratory Study*. *Journal AOAC*, 2013. 96, 1, 77-85.

OkaTest complies with the requirements established under chapter III A (4) a, b and c from Appendix III of the European Regulation (EC) 2074/2005 and can be used as complementary method.

For further information, please visit the European Reference Laboratory website:

http://aesan.mssi.gob.es/en/CRLMB/web/otros_procedimientos/other_crmb_standard_operating_procedures.shtml

OBJETIVO

Test para la determinación cuantitativa de Ácido Okadaico (OA) y otras toxinas del grupo del OA, incluyendo DTX1, DTX2 y DTX3. Consiste en un ensayo colorimétrico de inhibición de la actividad enzimática de una fosfatasa. Este método es aplicable a especies como mejillones, almejas, berberechos, vieiras, etc.

PRINCIPIO

Okatest es un test basado en la inhibición de la actividad enzimática de una fosfatasa (PP2A) por toxinas del grupo del ácido okadaico. En condiciones normales, la fosfatasa es capaz de hidrolizar un sustrato específico obteniéndose un producto que puede ser detectado a 405 nm. En presencia de toxina diarreica se producirá una inhibición de la actividad enzimática proporcional a la cantidad de toxina diarreica presente en la muestra. Mediante la utilización de una curva de calibrado se pueden obtener los valores de concentración de toxina presentes en la muestra analizada.

COMPONENTES DEL KIT

	Kit de 48 Tests	Kit de 96 Tests
Tiras de 8 pocillos de placa microtiter	6	12
Fosfatasa (<i>Phosphatase</i>)	2	4
Set de patrones de ácido okadaico (<i>Okadaic acid 0.5, 0.8, 1.2, 1.8 y 2.8 nM</i>)	1	1
Sustrato Cromogénico (<i>Chromogenic Substrate</i>)	1	1
Solución de Dilución de la Fosfatasa (<i>Phosphatase Dilution Buffer</i>)	1	1
Solución Tamponante (<i>Stock Buffer Solution</i>)	1	1
Solución Stop (<i>Stop Solution</i>)	1	1
Lámina adhesiva	1	2
Guión de instrucciones	1	1

MATERIAL Y REACTIVOS ADICIONALES NECESARIOS

- Micropipetas
- Homogeneizador (e.j. Ultraturax) o mortero
- Incubador a $30 \pm 2^\circ\text{C}$. (Ej. FX Incubator Ref ZE/FX, de ZEULAB)
- Lector de placas microtiter con filtro a 405 nm.
- Baño termostático $76 \pm 2^\circ\text{C}$
- Metanol (grado analítico)
- NaOH (grado analítico)
- HCl (grado analítico)
- Agua desionizada (al menos de grado 2, ISO 3696)
- Tubos de centrifuga de 50 mL
- Centrifuga
- Agitador para tubos (tipo vortex)

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SOLUCIONES

- 1.- Estándares de Ácido Okadaico: Es muy importante agitar bien estas disoluciones justo antes de su utilización (p.e.: en vortex), para asegurar su homogeneidad
- 2.- Sustrato Cromogénico: esta solución contiene una resina estabilizante que no debe añadirse a los pocillos. Con este fin, se recomienda transvasar el volumen a utilizar a un vial transparente (p.e.: eppendorf o tubo de ensayo), asegurándose de no coger resina, y de ahí pipetear a los pocillos. *Nota:* no usar esta solución si la absorbancia de 90 μ L es superior a 0.6.
- 3.- Preparación de la Fosfatasa: reconstituir el liofilizado de Fosfatasa (*Phosphatase*) en 2.0 mL de Solución de Dilución de la Fosfatasa (*Phosphatase Dilution Buffer*). Mantener la solución a temperatura ambiente ($22 \pm 2^{\circ}\text{C}$) y con agitación suave durante 1 hora para asegurar así la correcta hidratación del liofilizado. No usar el agitador de tubos en ningún momento. Una vez reconstituido el enzima, mantenerlo en condiciones de refrigeración. No conservar la solución de Fosfatasa para su uso en días posteriores. Cada vial de Fosfatasa contiene la cantidad necesaria para 24 pocillos. Si se va a utilizar más de uno, disolver cada vial como se ha explicado anteriormente y mezclar el contenido de todos en uno único antes de usar. Agitar suavemente antes de su utilización. **Atención:** el liofilizado posee una coloración azulada y al reconstituirlo se convierte en marrón. Si observa que este reactivo posee una coloración marrón antes de reconstituirlo, no usarlo, ya que podría estar dañado.
- 4.- Solución Tamponante x1: diluir la *Stock Buffer Solution* incluida en el kit, mezclando 1 volumen de esta solución con 9 volúmenes de agua desionizada. Preparar sólo la que se vaya a utilizar en el momento y mantener en refrigeración hasta entonces.
- 5.- NaOH 2.5 N: pesar 100 g de NaOH y disolver en 500 mL de agua desionizada. Seguidamente, enrasar hasta un volumen final de 1000 mL usando un matraz aforado.
- 6.- HCl 2.5 N: Añadir 205 mL de HCl (37 %) a 400 mL de agua desionizada. Mezclar y enrasar hasta 1000 mL con agua desionizada usando un matraz aforado.

EXTRACCIÓN DE LAS MUESTRAS

El método de preparación de muestras que se describe a continuación incluye una etapa de hidrólisis que permite la detección de todas las formas tóxicas de ácido okadaico (ácido okadaico y dinofisistoxinas).

- 1.- Limpiar la superficie externa del molusco con agua.
- 2.- Abrir los moluscos seccionando los músculos aductores.
- 3.- Lavar el contenido de las conchas con agua hasta conseguir eliminar todas las sustancias extrañas que puedan contener.
- 4.- Separar la carne de las conchas, retirando todos los músculos o tejidos que estén en contacto con ellas.
- 5.- Colocarlos en un papel de filtro y dejarlos secar durante unos minutos.
Se recomienda el uso de tubos calibrados para centrifuga de 50 mL durante las siguientes etapas de hidrólisis para evitar pérdidas por transvase de líquidos.
- 6.- Triturar el tejido hasta obtener una muestra homogénea, tomar 5 g (peso húmedo) y extraer con 25 mL de Metanol durante 2 minutos, usando un agitador para tubos.
- 7.- Centrifugar el homogeneizado a 2000 g durante 10 minutos a 4 $^{\circ}\text{C}$. Al sobrenadante lo llamaremos *extracto metanólico* y lo pasaremos a otro tubo de centrifuga por decantación.

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- 8.- Tomar 640 μL del *extracto metanólico* y transvasarlo a un tubo para centrifuga nuevo.
- 9.- Añadir 100 μL de NaOH 2.5 N.
- 10.- Cerrar y calentar la muestra a 76 ± 2 °C durante 40 minutos.
- 11.- Sin dejar enfriar, añadir 80 μL de HCl 2.5 N
- 12.- Añadir Solución Tamponante x1 hasta un volumen final de 20 mL.

PROCEDIMIENTO DE ENSAYO

Atención:

En este ensayo se usan reactivos en volúmenes pequeños y se debe tener especial cuidado cuando se añaden a la placa:

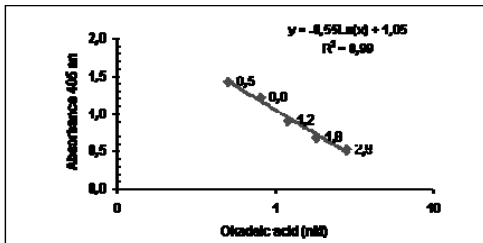
- Asegurarse de que las pipetas están calibradas antes de realizar el ensayo.
- Usar pipetas de 100 ó 200 μL de volumen máximo.
- Comprobar que la temperatura del incubador está estabilizada antes de su uso.

Es aconsejable aplicar las muestras y patrones por duplicado.

- 1.- Aplicar 50 μL de cada estándar o muestra.
- 2.- Aplicar en cada pocillo 70 μL de la Solución de Fosfatasa. Mezclar bien golpeando suavemente en el lateral de la placa.
- 3.- Tapar la placa con la lámina adhesiva incluida en el kit e incubar a 30 ± 2 °C durante 20 ± 0.5 minutos.
- 4.- Aplicar 90 μL en cada pocillo de Sustrato Cromogénico y tapar la placa con la lámina adhesiva.
- 5.- Incubar a 30 ± 2 °C durante 30 ± 0.5 minutos.
- 6.- Retirar la lámina adhesiva y añadir en cada pocillo 70 μL de Solución Stop.
- 7.- Leer la absorbancia a 405 nm en un lector de placas microtiter.

REPRESENTACIÓN Y CÁLCULO DE LOS RESULTADOS

- 1.- Obtener una curva de calibrado representando las absorbancias en el eje de ordenadas frente a las concentraciones de ácido okadaico en el eje de abscisas (este último en escala logarítmica). A continuación se muestra un ejemplo de curva patrón. R^2 deberá ser mayor o igual a 0.96.



- 2.- A partir de la curva de calibrado obtener los valores de ácido okadaico de las muestras (Cs) por interpolación o aplicando la ecuación correspondiente:

$$x = \text{EXP} (y - b/a)$$

x: concentración de ácido okadaico en la muestra

y: absorbancia de la muestra

*ZEULAB puede proporcionar una plantilla Excel para calcular los resultados. Para más información contacte con nosotros.

- 3.- Calcular la concentración de toxinas diarreicas en el tejido (Ct) a partir de la siguiente fórmula:

$$Ct (\mu\text{g}/\text{kg}) = \frac{Cs (\text{nM}) \times FD \times PM (\text{g}/\text{mol}) \times Ve (\text{L})}{Mt (\text{g})}$$

Ct: Concentración de toxinas en tejido; Cs: Concentración de toxinas de cada muestra aplicada en el pocillo; FD: Factor de dilución del extracto metanólico en la preparación de la muestra (p.e. 640 $\mu\text{L}/20 \text{ mL} \rightarrow \times 31.25$); PM: Peso molecular ácido okadaico = 805; Ve: Volumen de extracto metanólico obtenido (0.025L); Mt: Masa de tejido pesada inicialmente (5 g).

Ej.: Para una muestra 1.5 nM de OA: 1.5 nM \times 31.25 \times 805 g/mol \times 0.025 L / 5 g = 189 μg eq OA/kg

NOTA: Aquellas muestras cuya concentración (Cs) esté fuera del rango de trabajo (< 0.5 nM ó > 2.8 nM), los resultados se expresarán como < 0.5 nM (ó < 63 $\mu\text{g}/\text{Kg}$) ó > 2.8 nM (ó > 352 $\mu\text{g}/\text{kg}$) respectivamente.

Muestras con absorbancias inferiores a las obtenidas para el patrón 2.8 nM pueden ser analizadas de nuevo basándose en una dilución máxima de 1:4 del extracto metanólico.

ESTABILIDAD Y ALMACENAMIENTO

Conservar los componentes del kit de 4 -12 °C y en oscuridad. El kit tiene una estabilidad de 8 meses en las condiciones de conservación anteriormente indicadas.

SEGURIDAD

Se recomienda seguir unas prácticas correctas de laboratorio, así como el empleo de ropa y material de seguridad adecuados para el desarrollo del test. Evitar el contacto directo con la piel. No ingerir.

***Atención:** El ácido okadaico es un producto tóxico, para su manejo es imprescindible el uso de guantes y trabajar con precaución.

Puede solicitar la hoja de seguridad del producto contactando con su distribuidor habitual o fabricante.

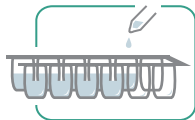
BIBLIOGRAFÍA

- 1.- Takai, A.; Bialojan, C.; Troschka, M.; Rüegg, J.C. *Smooth muscle myosin phosphatase inhibition and force enhancement by black sponge toxin. FEBS Lett.* 1987, 21781-21784.
- 2.- Smienk H., Calvo D., Razquin P., Domínguez E. & Mata L. *Single Laboratory Validation of A Ready-to-Use Phosphatase Inhibition Assay for Detection of Okadaic Acid Toxins. Toxins,* 2012, 5, 339-352.
- 3.- Smienk H., Domínguez E., Rodríguez-Velasco M.L. Clarke D., Katrin K., Katikou P., Cabado A.G., Otero A., Vieties J.M. Razquin P., and Mata L. *Quantitative Determination of the Okadaic Acid Toxins Group by a Colorimetric Phosphatase Inhibition Assay: Interlaboratory Study. Journal AOAC,* 2013. 96, 1, 77-85.

OkaTest cumple con los requisitos del capítulo III A (4) a, b y c del Anexo III de la Regulación Europea (EC) 2074/2005 y puede ser usado como método complementario tal y como indica el Laboratorio de Referencia Europeo en su página web: http://aesn.mssi.gob.es/en/CRLMB/web/otros_procedimientos/other_crlmb_standard_operating_procedures.shtml

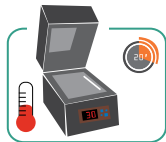
**FLOWCHART
PROCEDURE**

**ESQUEMA DEL
PROCEDIMIENTO**



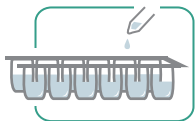
1. Add 50 μ L samples/standars
2. Add 70 μ L Phosphatase Solution

1. *Añadir 50 μ L muestras/estándares*
2. *Aplicar 70 μ L Solución de Fosfatasa*



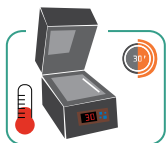
3. Incubate 20 min at 30°C

3. *Incubar 20 min a 30°C*



4. Add 90 μ L Cromogenic Substrate

4. *Añadir 90 μ L Sustrato Cromogénico*



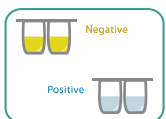
5. Incubate 30 min at 30°C

5. *Incubar 30 min a 30°C*



6. Add 70 μ L Stop Solution

6. *Añadir 70 μ L Solución Stop*



7. Read absorbance at 405 nm

7. *Leer absorbancia a 405 nm*

DSP PPIA kit-OkaTest

Single Laboratory Validation Report

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1- EXECUTIVE SUMMARY

The **DSP PPIA** (commercial name **OkaTest**) is a test for detection of Okadaic Acid (OA) and other carboxylic toxins of the OA group including DTX1, DTX2 and DTX3 by a colorimetric phosphatase inhibition assay. It is a rapid and simple method suitable for quantitative determination of the OA- toxins group from 63 to 352 µg of OA equivalents per Kg, including the maximum limit established as 160 µg of OA equivalents /Kg in the Commission Regulation of 29 April 2004 (Regulation (EC) 853/2004). Test applicable to shellfish species such as mussels, clams, oysters and scallops.

The **OkaTest** kit was developed by ZEULAB (previous name ZEU- INMUNOTEC) based on the research work carried out by Vieytes et al. The method uses the inhibitory activity of OA and DTXs against the enzyme phosphate, which is responsible for their toxic effect, for the detection of OA-toxins group in molluscs. **OkaTest** uses a colorimetric detection system (Takai and Mieskes, 1991), while the original method (Vieytes et al., 1997) was based on fluorimetric detection.

A single laboratory validation was carried out at ZEULAB, followed by a collaborative study with 16 laboratories from 11 different countries. Both validations have been published in scientific journals; Toxins in 2012 by Smienk et al. and Journal of AOAC in 2013 by Smienk et al., respectively. Besides, OkaTest participates annually in international proficiency exercises (Quasimeme, The Netherlands).

OkaTest complies with the requirements established by the European Regulation (EC) 2074/2005 as complementary to the reference method.

http://www.aecosan.msssi.gob.es/en/CRLMB/web/public_documents/seccion/other_crlmb_standard_operating_procedures.htm

Furthermore, **OkaTest** has been compared with other methodologies and using samples from the USA, UK and Argentina (Bich-Thuy et al., 2013, Turner & Goya, 2016 and Johnson et al., 2016).

This report shows the data obtained in the initial single laboratory validation (Smienk et al, 2012) that has been completed with additional information requested by the ISSC. Following a summary of the validation parameters:

Parameter	Results
Accuracy/Trueness	98,00%
Measurement Uncertainty	14.92 - 31.08 µg equivalentes OA /kg
Precision	
Repeatability:	1,4%- 3,9 % (Mean= 2,65%)
Reproducibility	0,8 %-17,7% (Mean= 6,45%)
Recovery	Okadaic acid: 78-114%
	DTX-1: 79-102%
	DTX-2: 83-94%
Working Range	63 - 352 µg equivalentes OA /kg
Limit of Detection (LOD)	44 µg equivalentes OA /kg
Limit of Quantification (LOQ)	56 µg equivalentes OA/kg

2- METHOD PRINCIPLE AND SCOPE

DSP PPIA (OkaTest) is a protein phosphatase inhibition assay (PPIA), where the phosphatase activity is inhibited by the OA-toxins group, responsible for diarrhetic shellfish poisoning (DSP). The PPIAs have been identified for a long time as an alternative for the detection of the OA-toxins, as Ser/Thr phosphatases are known to be their natural target (Bialojan & Takai, 1988). Under normal circumstances, a phosphatase enzyme is able to hydrolyse a specific substrate producing a reagent that can be detected by absorbance measurement (405 nm). Samples containing OA toxins will inhibit the enzyme activity proportionally to the amount of toxin contained in the sample.

OkaTest is applicable to shellfish species such as mussels, clams, oysters and scallops. It is a quantitative method for determination of the OA- toxins group, where concentration of toxins present in the sample is calculated using a standard curve.

OkaTest includes five OA standards (0.5, 0.8, 1.2, 1.8 and 2.8 nM), phosphatase enzyme and substrate reagents ready to use.

The test procedure is extensively described in the user manual G-COM-OA.06.

3- VALIDATION

To evaluate the performance of the OkaTest kit, accuracy, uncertainty, precision, limit of detection and quantification were calculated. The assay temperature, incubation times and other variables affecting ruggedness, together with specificity and matrix effects were also evaluated. Finally, a method comparison was carried out.

3.1 Accuracy/Trueness

To estimate the accuracy of the method 20 blank mussel samples (*Mytilus edulis*) were spiked with OA at 80, 120, 160, 240 and 300 µg/kg. Percentage of recoveries were calculated and are shown in Table 1.

Table 1. Recovery values from 20 different mussels samples spiked with OA at different levels along the working range. Mean (M), standard deviation (SD) and relative standard deviation (RSD). ND < 63 µg/kg

Theoretical Spike	µg OA equivalents/kg		Recovery	Mean	SD	RDS
	Before spiked	After spiked				
80	ND	73	91.3%			
80	ND	91	113.8%			
80	ND	87	108.8%	112.5%	0.18	16.68%
80	ND	112	140.0%			
80	ND	87	108.8%			
120	ND	133	110.8%	106.7%	0.06	5.52%
120	ND	123	102.5%			
160	ND	128	80.0%			
160	ND	169	105.6%	98.8%	0.13	12.98%
160	ND	173	108.1%			
160	ND	162	101.3%			
200	ND	186	93.0%			
200	ND	185	92.5%	91.3%	0.30	0.27%
200	ND	177	88.5			
240	ND	219	91.3%			
240	ND	205	85.4%	96.1%	0.21	21.59%
240	ND	195	81.3%			
240	66	304	126.7%			
300	ND	250	83.3%	82.7%	0.01	1.14%
300	ND	246	82.0%			

3.2. Measure of Uncertainty

Measurement of uncertainty was calculated using the results obtained in the accuracy experiment considering a confidence interval of 95%. Mean and standard deviation of the difference between the concentration of the spiked sample and the spiked amount were calculated. The coefficient of confidence (Z) and maximum error (E. max) were then determined (Table 2) according to the following equation:

$$E. \max = Z_{\alpha/2} * SD / \sqrt{n}, \text{ where}$$

E. max: maximum error, **Z:** confidence coefficient; **α** 95% confidence interval, **SD:** standard deviation, **n:** number of samples.

Table 2. Estimation of uncertainty based on recovery data from 20 different mussels. ABS: absolute value of differences between OA concentration in spiked samples and spike concentration. Z= coefficient of confidence. SD= standard deviation. ABS E. max=absolute value of maximum error. ND < 63 µg/kg

Sample	Spike (OA µg/kg)	Blank Sample µg OA equiv. /kg	Spiked Sample	Recovery	ABS differences	Mean	SD	ABS E. Max
1	80	ND	73	91.3%	7	13	10.87	9.53
2	80	ND	91	113.8%	11			
3	80	ND	87	108.8%	7			
4	80	ND	112	140.0%	32			
5	80	ND	87	108.8%	7			
6	120	ND	133	110.8%	13	8	7.07	6.20
7	120	ND	123	102.5%	3			
8	160	ND	128	80.0%	32			
9	160	ND	169	105.6%	9			
10	160	ND	173	108.1%	13			
11	160	ND	162	101.3%	2	14	12.83	11.25
12	200	ND	186	93.0%	14			
13	200	ND	185	92.5%	15			
14	200	ND	177	88.5%	23	17	4.95	4.34
15	240	ND	219	91.3%	21			
16	240	ND	205	85.4%	35			
17	240	ND	195	81.3%	45			
18	240	66	304	126.7%	64	41	18.08	15.85
19	300	ND	250	83.3%	50			
20	300	ND	246	82.0%	54	52	2.83	2.48
Mean					23			
SD					18.44			
ABS E. Max					8.08			

3.3. Precision

To determine the precision of the method, relative standards deviation (RSD) for repetibility and reproducibility were calculated.

To calculate repeatability eight replicates of two mussel samples at two levels of concentration were analysed on the same day. Mean, standard deviation and relative standard deviation were calculated. The RSD obtained for the samples tested were, 1.4 and 3.9%, respectively. These values are far below the reference value of 15% (Horwitz W., 2002).

Table 3. Repeatability of 2 different mussel samples. Mean, standard deviation (SD) and relative standard deviation (RSD).

Repetition	Sample 1 ($\mu\text{g OA equiv./kg}$)	Sample 2 ($\mu\text{g OA equiv./kg}$)
1	269	124
2	276	125
3	276	131
4	273	129
5	280	121
6	278	117
7	281	127
8	275	118
Mean	276	124
SD	3.9	4.8
RSD	1.4%	3.9%

Intermediate precision/Reproducibility

Intermediate precision was estimated by testing 13 different samples (10 mussel samples and 3 from other species) at different levels of concentration on 3 different days by the same analyst (Table 4).

Mean values, standard deviation and relative standard deviation were calculated. An average of 6.45% of RSD was calculated for all the samples with different levels of concentration. Only sample 3, at a concentration below the regulatory limit showed a RSD above 15%, which is the variability expected for this concentration range (Horwitz, 2002).

Table 4. Reproducibility of thirteen different mussel (*Mytilus edulis*), king scallop (*Pecten maximus*) and clam (*Venerupis pullastra* and *V. vomboides*) samples. Mean, standard deviation (SD), relative standard deviation (RSD) were calculated.

Sample	Matrix	Day 1	Day 2	Day 3	Mean	SD	RSD
		$\mu\text{ OA equivalents /kg}$					
1	Mussel	82	94	90	88	6.17	7.0%
2	Mussel	106	95	90	97	8.05	8.3%
3	Mussel	98	101	72	90	15.95	17.7%
4	Mussel	109	106	95	101	7.80	7.8%
5	King Scallop	125	108	117	117	8.20	7.0%
6	Mussel	122	132	113	122	9.57	7.8%
7	Mussel	196	196	215	202	10.57	5.2%
8	Mussel	211	227	187	208	19.84	9.5%
9	Clam	261	251	260	257	5.51	2.1%
10	Mussel	257	250	258	255	4.36	1.7%
11	Mussel	250	253	281	261	16.90	6.5%
12	Mussel	277	279	289	282	6.62	2.4%
13	Clam	285	285	281	284	2.31	0.8%

The intermediate precision was also further evaluated in a collaborative study with 5 samples analysed by 16 different laboratories. Values of 11.2% and 13.2% were determined as the

highest relative standard deviation for repeatability and reproducibility, respectively (Smienk et al 2013).

3.4. Recovery

Recovery was calculated by spiking mussel and scallop samples (*Mytilus edulis* and *Pecten maximus*, respectively) with okadaic acid (OA) at 0.5, 1 and 1.5 times the regulatory limit. Samples were also spiked with 80, 160 and 240 µg/kg of DTX-1 and 80 and 160 µg/kg of DTX-2. Three to five repetitions of each concentration were analysed on different days. Results are shown in Tables 5 and 6.

Table 5. Results (µg OA equivalents/kg) from recovery of OA in mussel and scallop samples at 80, 160 and 240 µg/kg. Standard deviation (SD), relative standard deviation (RSD) and recovery were calculated. ND= <63 µg/kg).

Repetition	Mussel				King Scallop			
	spiked OA (µg/kg)							
	0	80	160	240	0	80	160	240
1	86	158	230	271	ND	82	162	252
2	87	134	211	282	ND	84	142	218
3	87	178	216	257	ND	89	150	268
4	95	193	253	298	ND	102	177	268
5	95	191	257	280	ND	99	158	271
Mean	90	171	233	277	-	91	157	255
SD	4.8	25.0	20.9	15.1	-	9.0	13.3	22.2
RSD	5.4%	14.6%	8.9%	5.4%	-	9.9%	8.4%	8.7%
Recovery	-	101%	90%	78%	-	114%	98%	106%

Table 6. Results (µg OA equivalents/kg) from recovery of DTX-1 and DTX-2 in mussel and scallop samples spiked at 80, 160 and 240 µg/kg. Mean, Standard deviation (SD), relative standard deviation (RSD) and recovery were calculated. ND= <63 µg/kg).

Repetition	King scallop						Mussel			
	spiked DTX - 1 (µg/kg)						spiked DTX2 (µg/kg)			
	0	80	160	240	0	160	0	80	0	160
1	ND	63	101	211	ND	145	86	157	ND	128
2	ND	91	127	179	ND	156	101	163	ND	130
3	ND	81	132	175	ND	151	-	-	ND	124
4	ND	82	132	261	-	-	-	-	-	-
5	ND	93	140	228	-	-	-	-	-	-
Mean	ND	82	126	211	ND	151	93.5	160	ND	127
SD	-	11.9	14.8	35.6	-	5.5	-	4.2	-	2.7
RSDr	-	14.5%	11.7%	16.9%	-	3.7%	-	2.7%	-	2.1%
Recovery	-	102%	79%	88%	-	94%	-	83%	-	80%

The mean of recoveries obtained for the different concentrations tested and toxins were acceptable and ranged from 78 to 114%.

3.5. Specificity

Specificity was studied by determining the possible interferences caused by other lipophilic toxins such as Azaspirazides (AZA), Yessotoxins (YTX) and Pectenotoxins (PTX).

A mussel sample naturally contaminated was spiked, on two different days, with 160 µg/kg of AZA-1 (NRC, Institute for Marine Biosciences, Canada), 160 µg/kg PTX-2 (Cifga laboratories, Spain) and 1000 µg/kg of YTX (NRC, Institute for Marine Biosciences, Canada) and concentration of OA determined following the kits' instructions. Results obtained for spiked and non-spiked samples were very similar and within the method variability, showing no interferences by the toxins tested.

Table 6. Results obtained from spiking a mussel sample with 160 µg/kg of azaspirazides (AZA), 160 µg/kg of pectenotoxins (PTX) and 1000 µg/kg of yessotoxins (YTX).

Spiked Mussel	Day 1 µg equiv. OA /kg	Day 2 µg equiv. OA /kg
0	82	82
160 (µg/kg) PTX-2	83	79
160 (µg/kg) AZA-1	82	73
1000 (µg/kg) YTX	82	82

3.6. Working Range and Linear Ranges

The working range is understood as the range of OA concentrations that do correctly adapt to the fitting procedure. The working range of the assay depends on the quantity and quality of the phosphatase present. Therefore, assays were performed with at least 3 different phosphatase batches and the "goodness of fit" was evaluated according to the kits' specifications ($R^2 > 0.96$) with standard concentrations rising from 0.25 to 3.5 nM OA.

Figure 1 shows the results of three assays covering the range from 0.25 to 3.0 nM OA as this was the range that always fitted correctly ($R^2 > 0.96$). This covers sufficiently the actual range of the standards in the kit (0.5 to 2.8 nM OA).

The linearity of an assay was tested to find out whether the response of this assay is a function of the concentration of the analyte. The OkaTest assay uses a logarithmic fitting procedure.

As such the linearity of the assays' response was tested by 'backcalculation' of the standard concentration. For 'backcalculation' the equation of the standard curves used to calculate the concentration of these standards from their absorbances (Table 7).

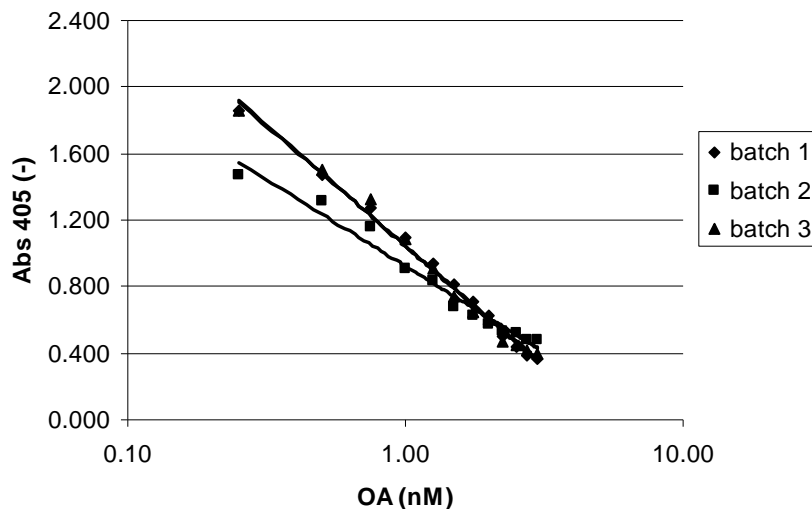


Fig 1. Working range of the assay for 3 different phosphatase batches. R^2 : 0.99, 0.98 and 0.99 for batch 1, batch 2 and batch 3, respectively. Working range from 0.25 to 3.0 nM OA.

Table 7. Linearity of the assay. OA (nM) was calculated by using the standard curve of batch 1.

Standards OA (nM)	Batch 1 OA (nM)	Batch 2 OA (nM)
0.5	0.6	0.5
0.8	0.7	0.7
1.2	1.1	1.2
1.8	1.9	1.9
2.8	2.9	2.8

To check the linearity of the response, the theoretical concentration was compared to the calculated concentration for both batches (see figure 2 for the results shown in Table 1) and a linear fit was performed. The Pearson correlation coefficient (R^2) for batch 1 was 0.99 and 1.00 for batch 2.

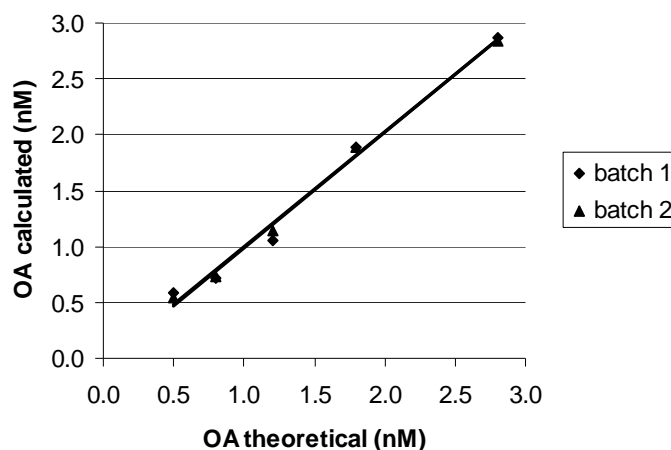


Fig 2. Comparison of the theoretical and calculated standard concentration. Concentration were 'backcalculated' by using the standard curve obtained with batch 1. R^2 : 0.99 and 1.00 for batches 1 and 2, respectively.

The linearity was also determined by testing 10 blank mussel samples spiked at 80, 160, 200, 240 and 300 $\mu\text{g}/\text{kg}$ (Table 8). OA concentrations obtained were divided by the spiked concentration (relative recovery). Mean of relative recovery per concentration was plot against the spiked concentration and curve equation to observe the relative response (Figure 3).

Table 8. Assay linearity. Results obtained from 10 blank samples spiked with 80, 120, 160, 200 and 240 $\mu\text{g}/\text{kg}$ to determine linearity of the assay.

Spike OA $\mu\text{g}/\text{kg}$	After spiked μg OA equiv. /kg	Relative recovery	Mean Relative Recovery
80	91	1,14	1,11
80	87	1,09	
160	169	1,06	0,96
160	162	1,01	
200	186	0,93	0,91
200	177	0,88	
240	219	0,91	0,88
240	205	0,85	
300	250	0,83	0,83
300	246	0,82	

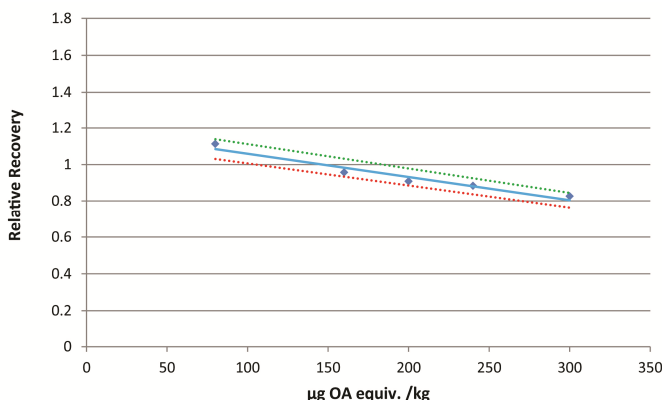


Figure 3. Assay linearity. Relative recovery data plot against spiked OA concentration µg/kg in solid blue line. Green and red dotted lines were obtained by multiplying the OA concentration by 0.95 and 1.05.

3.7. Limit of detection and Limit of quantification

To estimate the LOD and LOQ a blank mussel material was extracted ten times and analyzed according to the kits’ instructions. The mean and standard deviation were calculated and the limit of detection was estimated by the equation below:

$$LOD_{99\%} = X + 3SD$$

The LOQ (the lowest concentration that can be determined with an acceptable level of repeatability precision and trueness) was estimated using the same data and equation, but applying a higher factor:

$$LOQ_{99\%} = X + 10SD$$

The mean result obtained for the blank sample was 38 µg/kg. The estimated LOD and LOQ were 44 µg/kg and 56 µg/kg, respectively (Table 11).

Table 9. Quantification of the standard solvent (10 repetitions) as OA concentration equivalents (µg/kg) to estimate the LOD and LOQ. Mean, standard deviation (SD) and relative standard deviation (RSD).

Repetition	µg OA equivalents /kg
1	36
2	38
3	36
4	37
5	41
6	37
7	40
8	38
9	40
10	38
mean	38
SD	1.8
RSD	4.6%
LOD	44
LOQ	56

3.8. Ruggedness

The influence of different experimental conditions critical for the kits' performance such as assay temperature, incubation times or reaction component volumes were evaluated. The ruggedness between batches with spiked mussel samples was also evaluated.

3.8.1- Assay temperature

The hydrolysis of the substrate by the phosphatase is temperature dependent and shows the typical behaviour of an enzymatic reaction with higher reaction rates close to the optimum temperature (37°C). However, a lower assay temperature was chosen to guarantee enzyme stability during the assay and to get stable reaction rates. The assay was tested at temperatures varying from 20 to 40 °C. 30 °C was chosen as the optimum temperature. At this temperature a 2 °C variation can be expected in any incubator. So, to show the influence of this temperature variation, 3 samples were quantified performing a complete assay (standard curve and samples) at each of these temperatures (Table 10).

Table 10. Influence of the assay temperature on the results of the test. The mean, standard deviation (SD) and relative standard deviation (RSD) were calculated.

Sample	28 °C	30 °C	32 °C	mean	SD	RSD
1	104	100	97	100	3.4	3.4%
2	176	173	176	175	1.7	1.0%
3	302	303	298	301	2.6	0.9%

Mean and relative standard deviation were calculated. For all three samples RSD were below the 15%, variation that can be expected at this concentration (Horwitz, 2002).

3.8.2- Assay incubation times

The assay consists of two different incubation steps that could affect the outcome of the test. During the first incubation the sample and the phosphatase are mixed, and the inhibition reaction should reach its endpoint. Following, the substrate is added and the plates are incubated for the second time. The main risk of this incubation step is phosphatase activity loss.

To determine the influence of time on the first incubation of the assay (normally 20 minutes), this step was varied between 18 and 24 min, while maintaining the rest of the assays' conditions according the kits' instructions. Three control samples were quantified and the variation in the relative standard deviation was evaluated. For each of the incubation conditions an independent assay was performed (Table 11).

Table 11. Influence of time on the first incubation of the assay. The mean, standard deviation (SD) and relative standard deviation (RSD) were calculated.

Sample	- 2 min	0	+ 2 min	+ 4 min	mean	SD	RSD
1	85	87	87	90	88	2.1	2.4%
2	152	155	161	164	158	5.7	3.6%
3	311	291	317	320	310	12.9	4.2%

In all cases the assay complied with the criterion ($R^2 > 0.96$). The relative standard deviations were comparable to those obtained when performing the test under standard conditions (highest 4.2%).

The second incubation was evaluated similarly. A 10% error from normal incubation time (30 minutes) was applied, adding some extra time (up to 20% or 6 minutes). The assay was performed as described; although no stopping solution was added to permit reading the same assay. The RSD was 2.9% at highest, a bit lower than the ones obtained for the first incubation time (Table 11).

Table 12. Influence of the incubation time (2nd incubation with the substrate) on the assay. The mean, standard deviation (SD) and relative standard deviation (RSD) were calculated.

Sample	- 3 min	0	+ 3 min	+ 6 min	mean	SD	RSD
1	89	90	89	91	90	1.0	1.1%
2	143	152	145	149	147	4.3	2.9%
3	309	321	315	313	315	5.2	1.7%

3.8.3- Influence of pipetting volumes

The OkaTest assay consists of three pipetting steps of relatively small volumes. First, 50 μ L samples of standards are applied in duplicate and 70 μ L of phosphatase is added. Then, after the first incubation, 80 μ L of substrate and finally 70 μ L of stopping solution are added. The influence of pipetting error was evaluated by introducing a 2 μ L systematic error in each of the pipetting steps, e.g. a -2 μ L error means pipetting 48, 68, 78 and 68 μ L for samples/standard, phosphatase, substrate and stopping solution, respectively. This relatively big error (4% of the sample volume) is quite above the systematic error that can be expected in correctly calibrated pipettes (2%), but it was chosen in order to get clear results for obvious interpretation. The RSD and error were evaluated (Table 13).

Table 13. Effect of the systematic pipetting error on the results of the test. Mean, standard deviation (SD), relative standard deviation (RSD) and error (Errormax) were calculated.

Sample	- 2 μL	0	+ 2 μL	mean	SD	RSD	E. Max*
1	83	85	93	87	4.9	5.6%	8.0 (9.4%)
2	161	148	156	155	6.7	4.3%	13 (8.8%)
3	303	289	304	299	8.5	2.8%	15 (5.1%)

*E. max = maximum difference from standard (0) conditions in $\mu\text{g}/\text{kg}$ and percentage.

The RSD was at highest 5.6% and in accordance with the values normally obtained with OkaTest. The error introduced changed from 9.4 to 5.1% of the standard conditions.

The effect of a single pipetting error was evaluated by introducing a 5 μL error in one of the pipetting steps. In this case, the standard curve was performed according the kits' instructions and the error was introduced in the samples that were quantified. For example, a -5 μL error in the phosphatase means that 65 μL phosphatase was added to 50 μL sample (in duplicate) after which the assay was performed as usual. Also in this case, a relatively big error was chosen (10-6.3 % error, depending on the assay volume) (Table 14).

Table 14. Effect of a single pipetting error on the results of the test. Mean, standard deviation (SD), relative standard deviation (RSD) and error (Errormax) were calculated.

Variable	-5 μL	0	+ 5 μL	mean	SD	RSD	E Max*
Sample	132	148	173	151	20.5	13.6%	25 (17%)
Phosphatase	180	148	130	153	25.2	16.5%	32 (22%)
Substrate	167	148	159	158	9.6	6.1%	19 (13%)
Stop solution	170	148	153	157	11.6	7.4%	22 (15%)

*E. max = maximum difference from standard (0) conditions in $\mu\text{g}/\text{kg}$ and percentage.

Table 14 shows that pipetting errors in sample and phosphatase volume have the biggest effect and special care have to be taken when applying these. Also the logical tendencies can be seen; when applying less samples underestimation can be expected, while with the phosphatase occurs the contrary. This is to be expected, less phosphatase means more inhibitor per amount of phosphatase and so higher estimates of the toxin concentration. Table 14 also shows that high RSDr values (above 10%, ZEULAB in-house 5%) are a good indication for pipetting error. Substrate and stop solution pipetting errors seem to be much less important RSDr < 10%.

3.8.4- Influence of phosphatase solubility

In the previous paragraph was shown that the amount of phosphatase added to each well is important for correct quantification. The phosphatase is the only component of the kit that is not ready to use. It has to be dissolved previously and insufficient solubilisation could lead to

overestimation of the toxin concentration. Therefore the solubilisation time was evaluated by dissolving three phosphatase vials of the same batch for 30, 60 and 90 minutes (normal resuspension time use is 60 minutes), and always under agitation. Three control samples were quantified and the RSD was evaluated (table 15).

Table 15. Test results after dissolving the phosphatase for 30, the normal 60 and 90 minutes. The remaining part of the assay was performed according to the kits instructions. Mean, standard deviation (SD), relative standard deviation (RSD) were calculated.

Sample	30 min	60 min	90 min	Mean	SD	RSD
1	100	95	99	98	2.5	2.5%
2	167	151	157	158	8.0	5.0%
3	317	304	318	313	8.1	2.6%

The RSD values obtained were at highest 5.0% and comparable to those obtained for within batch variability (see table 15).

3.8.5- Ruggedness between batches in samples

The ruggedness of the assay with molluscs samples was also determined. Ten blank mussel samples were spiked at 80, 120, 160, 200 and 240 µg/kg of okadaic acid and tested following the kits' instructions in two different days and using two different batches. Differences between concentrations obtained in each batch for the different samples were calculated. Mean and standard deviation of the differences together with the experimental t-score and critical t values were also determined (Table 16).

$$t_{exp} = \frac{|Mean|}{\frac{s}{\sqrt{n}}}$$

Mean ≡ mean of the difference of skewness

s ≡ Standard deviation; n ≡ number of samples

The critical value was calculated for a significance of $\alpha = 0.05$ (95% confidence) for n-1 degrees of freedom. If the calculated value of experimental-t is less than the critical-t, we can affirm that the hypothesis is true, so that there is an equivalence between both methods.

The experimental t-score was smaller than the critical t-value ($t_{exp} < t_{crit}$; $1.42 < 2.26$); and so the range of skewness was acceptable. There is not significant difference between batch 1 samples and batch 2 concentrations.

Table 16. Results from testing 10 different mussel samples spiked at different concentrations and tested with two different batches in two different days. Mean, standard deviation (SD), experimental-t and critical-t values were calculated.

Sample	Batch No. 1	Batch No. 2	Differences between batches
	OA equivalents µg/kg		
1	91	74	-17
2	87	79	-8
3	133	102	-31
4	123	150	27
5	169	145	-24
6	162	177	15
7	186	177	-9
8	185	168	-17
9	219	174	-45
10	159	169	10
	Mean		-9.9
	SD		22.01
	Experimental t-score		1.42
	Critical t-value		2.26

The data was also analyzed using a Welch's test or unequal variances t-test, which is a two-sample location test used to check the hypothesis that two populations have equal means (H_0).

Therefore, considering that the null hypothesis (H_0) refers to the fact that the two batches do not show differences in the analysis of samples spiked with a known concentration of okadaic acid. Mean, variance and p-value were calculated (Table 17).

P-value (0.603) was higher than 0.05 ($0.603 > 0.05$), therefore we do not reject the null hypothesis. The observed difference between the sample's means is not convincing enough to say that the average value between both batches differing significantly.

Table 17. Mean, variance and p-value calculated for results obtained from 10 spiked mussel samples tested with bath 1 and batch 2 of OkaTest (results from Table 16).

	Batch 1	Batch 2
Mean	151.40	141.50
Variance	1812.93	1682.50
p value	0.603	

Residual values analysis evaluates the goodness of the test. A linear relationship is confirmed when the residues have symmetry around zero and a homogeneous random dispersion. Graphical representation is the most common methodology, being a very visual and simple method to evaluate symmetry. Residual standard values were also calculated (table 18) and

the distribution plotted. The adjustment is adequate since the residual values have a random and homogeneous distribution around 0, being between ± 2 (Figure 4).

Table 18. Residual standard values obtained for OA concentration results obtained for 10 mussel samples analysed with two different batches of OkaTest.

Sample	Residual standard values
1	5.51E-05
2	-0.39
3	0.79
4	-1.67
5	0.67
6	-0.98
7	0.13
8	0.46
9	1.78
10	-0.79

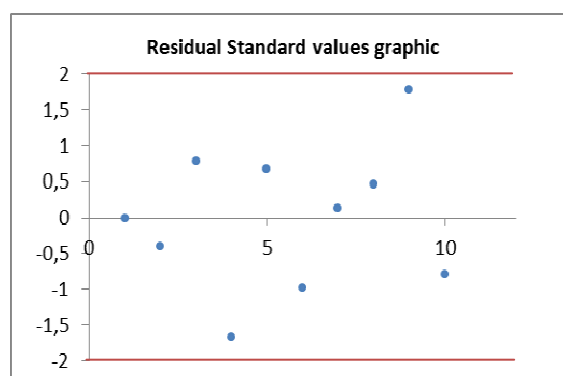


Figure 4. Distribution of residual standard values obtained for OA concentration results obtained for 10 mussel samples analysed with two different batches of OkaTest

3.9- Matrix Effects:

To determine the matrix effect 10 different molluscs' samples were tested according to the kit instructions and further diluted; where the final concentration of diluted samples was calculated multiplying by the appropriate dilution factor. Mean and SD of the differences between concentrations for diluted samples were calculated.

To evaluate if the concentrations obtained for diluted samples were within the assay variability and not due to matrix effect the experimental t-score and t-critical values were calculated (Table 19):

As the experimental t-score is smaller than the critical t-value ($0.93 < 2.26$) the skewness obtained is acceptable and does not indicate matrix effect.

Table 19. OA equivalents $\mu\text{g}/\text{kg}$ for 10 mussel samples tested a two different dilutions. Mean, standard deviation (SD), experimental-t and critical-t values were calculated.

Sample	Dilution 1	Dilution 2	Differences
	OA equiv. $\mu\text{g}/\text{kg}$		
1	40	37	-3
2	980	974	-6
3	29	30	1
4	620	628	8
5	595	560	-35
6	138	104	-34
7	1192	1287	95
8	1149	1318	169
9	118	108	-10
10	85	87	2
		mean	18.7
		SD	63.84
		Experimental- t score	0.93
		Critical-t value	2.26

3.10. Method comparison

A method comparison was performed with the mouse bioassay (MBA), reference method in Europe until 2011 and LC-MS/MS (current official reference method in Europe).

To compare results from OkaTest and MBA, values obtained by OkaTest with a concentration $\geq 160 \mu\text{g}/\text{kg}$ were regarded as positive while samples with a concentration $< 160 \mu\text{g}/\text{kg}$ were reported negative.

Twenty-three out of thirty-one samples tested positive for both methods and five samples were negative for both methods. However, three samples were positive for MBA and negative for OkaTest (Table 20). In all three samples OA toxins were detected, but below the regulatory limit of $160 \mu\text{g}/\text{kg}$ (144, 135 and $124 \mu\text{g}/\text{kg}$ OA toxins, respectively). Those samples were also tested by LC-MS/MS where two out of three results were above the regulatory limit as well by MBA, showing slightly higher quantifications compared to OkaTest (185, 152 and $177 \mu\text{g}/\text{kg}$ OA toxins $\mu\text{g}/\text{kg}$, respectively).

OkaTest was compared with LC-MS/MS for a total of 69 samples, where results from the reference method came from two different laboratories (Tables 20 and 21).

Table 20. Results from MBA, OkaTest and LC-MS/MS. Positive results (+): $\geq 160 \mu\text{g}/\text{kg}$. Negative result (-): $< 160 \mu\text{g}/\text{kg}$. *HPLC-MS results were not with toxicity factors. However only 4 samples contained DTX-2. LC-MS/MS carried out at the Reference Laboratory in Vigo.

Sample	Matrix	MBA	OkaTest	OkaTest $\mu\text{g OA equiv. /kg}$	LC-MS/MS $\mu\text{g OA /kg}$
1	Mussel	-	-	122	ND
2	Scallop	-	-	ND	ND
3	Mussel	-	-	ND	ND
4	Donax	-	-	97	82
5	Cockle	-	-	ND	ND
6	Mussel	+	+	196	158
7	Mussel	+	+	232	502
8	Mussel	+	+	268	ND
9	Scallop	+	+	264	184
10	Mussel	+	+	250	177
11	Mussel	+	+	265	288
12	Mussel	+	+	196	318
13	Mussel	+	+	>377	604
14	Mussel	+	+	>377	894
15	Mussel	+	+	277	390
16	Mussel	+	+	305	658
17	Mussel	+	+	306	414
18	Mussel	+	+	310	392
19	Mussel	+	+	>377	444
20	Mussel	+	+	315	329
21	Mussel	+	+	270	232
22	Mussel	+	+	277	235
23	Mussel	+	-	135	152
24	Mussel	+	+	164	98
25	Mussel	+	+	211	168
26	Mussel	+	+	251	209
27	Mussel	+	+	191	113
28	Mussel	+	-	124	177
29	Cockle	+	+	252	193
30	Mussel	+	+	216	247
31	Mussel	+	-	144	185
32	Mussel		-	ND	ND
33	Mussel		+	>377	357
34	Mussel		-	ND	292
35	Mussel		-	ND	ND
36	Mussel		-	ND	ND
37	Mussel		+	304	316

A comparison of OkaTest and the reference method LC-MS/MS was made for those samples which showed a quantitative value with both methods. The samples were analyzed by paired t-test to determine the equivalence of the two analytical methods, comparing both means to determine if the difference between the expected means surpasses the one produced randomly.



The hypothetical difference of Means should be zero (Null hypothesis H_0), which means that both methods are considered equivalents.

Table 21. Analysis t Student match pairs from results OkaTest and LC-MS/MS results from table 20.

	OkaTest	LC-MS
Mean	240.33	281.71
t-statistic	1.74	
P(T≤t) value (probability value) for the t-statistic (one-tailed)	0.048	
Critical value of a t-distribution (one-tailed)	1.72	
P(T≤t) value (probability value) for the t-statistic (two-tailed)	0.097	
Critical value of a t-distribution (two-tailed)	2.09	

The null hypothesis was accepted because critical- t two-tail < t Stat < t Critical two-tail (-2.09 < -1.74 < 2.09) and p (0.097)>0.05. The observed difference between the sample means (240.33 and 281.71) was not convincing enough to say that the average value between LC-MS and Okatest differ significantly.

Besides, the test t was applied manually to the difference of values obtained for each sample. For this application, the value of the experimental t-score statistic was calculated, as well as the critical t- value:

We could affirm that the hypothesis is true because the calculated experimental-t value was smaller than the critical-t value (1.65<2.08). The skewness is acceptable and the methods Okatest and LC-MS/MS are considered to be similar (Table 22).

Table 22. Mean, standard deviation (SD), experimental-t and critical-t values were calculated.

Mean	37.77
SD	107.56
Number of samples	22
Experimental t-score	1.65
Critical-t value	2.08

Table 23. Results from OkaTest and LC-MS/MS (EU harmonized SOP, v2, 2010)
Assays carried out by Jonathan Deeds from the FDA, US

Shellfish/Location	OkaTest	$\mu\text{g equiv. OA/Kg}$	LC-MS/MS		
	$\mu\text{g equiv. OA/Kg}$		OA	DTX1	DTX2
Softshell Clams (<i>Mya arenaria</i>) State: New York US East Coast Atlantic Ocean	336	373	255	118	ND
	315	325	202	105	ND
	295	307	217	108	ND
	285	260	136	69	ND
	240	205	171	89	ND
	190	155	102	53	ND
	118	75	49	26	ND
	<63	39	26	13	ND
	<63	ND	ND	ND	ND
Oysters (<i>Crassostrea virginica</i>) State: Texas US Gulf Coast Gulf of Mexico	322	563	563	ND	ND
	300	519	519	ND	ND
	245	202	202	ND	ND
	240	194	194	ND	ND
	239	221	221	ND	ND
	235	189	189	ND	ND
	198	189	189	ND	ND
	155	88	88	ND	ND
154	97	97	ND	ND	
	88	38	38	ND	ND
	<63	16	16	ND	ND
Mussels (<i>Mytilus edulis</i>) State: Washington US West Coast Pacific Ocean	>352	525	ND	525	ND
	266	272	ND	272	ND
	256	263	ND	263	ND
	171	165	ND	165	ND
	157	164	ND	164	ND
	141	131	ND	131	ND
	134	128	ND	128	ND
	127	121	ND	121	ND
	90	76	ND	76	ND
	<63	76	ND	76	ND
<63	33	ND	33	ND	
<63	ND	ND	ND	ND	

Mussels' results were statistically analyzed by applying a t-Student match pairs test to the results above the limit of quantification for each method.

The null hypothesis was accepted because the critical-t two-tail $< t \text{ Stat} < \text{Critical-t two-tail}$ ($-2.37 < 0.94 < 2.37$) and $p (0.8) > 0.05$. Therefore, we do not reject the null hypothesis. The observed difference between the sample means (167.75 and 165.00) is not convincing enough to say that the average value between LC-MS and Okatest differ significantly (Table 24).

Table 24. Analysis t Student match pairs from results OkaTest and LC-MS/MS results from table 22:

	OkaTest	LC-MS
Mean	167.75	165.00
t-statistic	0.94	
P(T≤t) value (probability value) for the t-statistic (one-tailed)	0.19	
Critical value of a t-distribution (one-tailed)	1.89	
P(T≤t) value (probability value) for the t-statistic (two-tailed)	0.38	
Critical value of a t-distribution (two-tailed)	2.36	

We applied the test t manually to the difference of values obtained for each sample. For this application the value of the experimental t-score statistic was calculated, as well as the critical-t value (Table 25).

Table 25. Mean, standard deviation (SD), experimental-t and critical-t values were calculated.

Mean	-2.75
SD	8.26
Number of samples	8
Experimental t-score	0.94
Critical-t value	2.36

We could affirm that the hypothesis is true because the calculated value of experimental-t is smaller than the critical-t ($0.94 < 2.36$). The skewness is acceptable and the values obtained by Okatest and LC-MS/MS are considered similar (Table 23).

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*Article***Single Laboratory Validation of A Ready-to-Use Phosphatase Inhibition Assay for Detection of Okadaic Acid Toxins****Henry G. F. Smienk, Dolores Calvo, Pedro Razquin, Elena Domínguez * and Luis Mata**

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Abstract: A phosphatase inhibition assay for detection of okadaic acid (OA) toxins in shellfish, OkaTest, was single laboratory validated according to international recognized guidelines (AOAC, EURACHEM). Special emphasis was placed on the ruggedness of the method and stability of the components. All reagents were stable for more than 6 months and the method was highly robust under normal laboratory conditions. The limit of detection and quantification were 44 and 56 µg/kg, respectively; both below the European legal limit of 160 µg/kg. The repeatability was evaluated with 2 naturally contaminated samples. The relative standard deviation (RSD) calculated was 1.4% at a level of 276 µg/kg and 3.9% at 124 µg/kg. Intermediate precision was estimated by testing 10 different samples (mussel and scallop) on three different days and ranged between 2.4 and 9.5%. The IC₅₀ values of the phosphatase used in this assay were determined for OA (1.2 nM), DTX-1 (1.6 nM) and DTX-2 (1.2 nM). The accuracy of the method was estimated by recovery testing for OA (mussel, 78–101%; king scallop, 98–114%), DTX-1 (king scallop, 79–102%) and DTX-2 (king scallop, 93%). Finally, the method was qualitatively compared to the mouse bioassay and LC-MS/MS.

Keywords: protein phosphatase inhibition assay (PPIA); protein phosphatase 2A (PP2A); validation; okadaic acid (OA); diarrheic shellfish poisoning (DSP)

1. Introduction

Diarrhetic shellfish poisoning (DSP) is a consequence of the ingestion of a series of lipophilic toxins produced by dinoflagellates that can be present in shellfish for human consumption. These lipophilic toxins can be subdivided into four groups: the okadaic acid group (OA-toxins) including the dinophysistoxins (DTX), the pectenotoxin group (PTX), the yessotoxin group (YTX) and finally the azaspiracids (AZA). Only the OA-toxins and AZA are known to cause gastrointestinal problems [1,2]. For many years the mouse bioassay (MBA) has been the official method of detection for lipophilic toxins in the European Union [3], but with the publication of Commission Regulation (EU) No. 15/2011 [4], LC-MS/MS has become the reference method for their determination. This regulation also states that alternative or complementary methods can be used as long as an equivalent level of public health protection is provided, and the method performance criteria stipulated by the European Union Reference Laboratory on Marine Biotoxins (EU-RLMB) are fulfilled. Such methods should be intra-laboratory validated and successfully tested under a recognized proficiency test scheme.

Protein phosphatase inhibition assays (PPIA) have been identified for a long time as an alternative for the detection of OA-toxins, as ser/thr phosphatases are known to be their natural target [5,6]. As such, a validated phosphatase inhibition assay can be very useful in lipophilic toxin detection, complementary to the more complex, expensive and time consuming LC-MS/MS; or as an alternative when only OA-toxins are present in the samples. Different laboratories have developed in-house PPIA with good qualifications, using colorimetric or fluorimetric substrates to monitor enzyme inhibition. [7–12]. A collaborative study was also performed with a fluorimetric assay [13]. However, specific equipment, not often available in routine testing laboratories, makes difficult the use of fluorimetric assays for monitoring purposes. Besides, fluorimetric substrates are less stable than colorimetric ones and therefore less appropriate for ready-to-use kits. A standardized commercial test based on PPIA has not been available until recently. In this paper, we present a single laboratory validation of a commercial colorimetric PP2A assay (OkaTest) for the determination of OA-toxins in bivalve mollusks.

2. Materials and Methods

2.1. Reagents and Equipment

OkaTest kit (formerly Toxiline-DSP): The kit includes a 96-well microtiter plate, four vials of lyophilized protein phosphatase 2A (PP2A), purified from human red blood cells, five OA standards (0.5, 0.8, 1.2, 1.8 and 2.8 nM) prepared from the OA Certified Reference Material (NRC CRM-OA-c, NRC-CNRC, Institute for Marine Biosciences), a liquid chromogenic substrate (p-Nitrophenyl phosphate), phosphatase dilution buffer and buffer solution.

Other reagents not included in the OkaTest kit: Methanol (Reagent grade, Carlo Erba), HCl (Reagent grade, 37% v/v, Carlo Erba), NaOH (Reagent grade, Scharlau), de-ionized water (type II, ISO 3696), certified Reference Materials (NRC CRM-DSP-MUS-b, NRC CRM-OA-c, NRC-CNRC, Institute for Marine Biosciences), DTX-1 (042-28661, Wako) and DTX2 (00-DTX2, Cifga).

Equipment: Ultra homogenizer (IKA werken), a water bath at 76 ± 2 °C (Raypa), a FX-incubator at 30 ± 2 °C (ZEU-INMUNOTEC), a microplate absorbance reader (405 nm \pm 10 nm wavelength

filter, Multiskan RC, Thermo-Labsystems), roller mixer, centrifuge, micropipettes, graduated 50 mL centrifuge tubes and laboratory glassware.

2.2. Sample Preparation

Market samples were thoroughly washed, the whole mollusk tissue recovered from the shell, and then blended. Portions of 5 ± 0.1 g were prepared and used for fresh testing, or stored frozen (below -15 °C) for future analysis. The portions were extracted by adding 25 mL of methanol (100% v/v) and mixing with a vortex for 2 min. The methanolic extract was separated by centrifugation for 10 min. at $2000 \times g$. To perform the hydrolysis, 640 μ L of the methanolic extract and 100 μ L of 3 N NaOH were mixed and incubated for 40 ± 1 min. at 76 ± 1 °C. To stop the reaction, 80 μ L of HCl were added and sample preparation buffer used to make up a final volume of 20 mL. For non-hydrolyzed samples, 640 μ L of methanolic extract were diluted up to 20 mL with sample preparation buffer. Hydrolysis was carried out in most samples unless otherwise specified.

2.3. Assay Procedure

The phosphatase solution was prepared by adding 2 mL of dilution buffer to each vial of lyophilized PP2A. To assure full hydration of the lyophilized enzyme, it was mixed gently for $1 \text{ h} \pm 5$ min. at room temperature (22 °C ± 2 °C) on a roller mixer. Then, 50 μ L of samples or ready-to-use OA standards (0.5, 0.8, 1.2, 1.8 and 2.8 nM), and 70 μ L of the prepared phosphatase solution were added in duplicate to a microwell plate. This mixture was equilibrated in an incubator for 20 ± 2 min. at 30 °C. Finally, 90 μ L of the chromogenic substrate were added to each well and incubated for 30 ± 2 min. at 30 °C. The absorbance was read at 405 nm.

2.4. Calculations

The results were calculated from a standard curve by plotting the absorbance values in a linear *y axis* and the concentration of OA in a logarithmic *x axis*, and using a logarithmic fitting. As an acceptability criterion for the assay, the Pearson correlation coefficient r^2 had to be greater than or equal to 0.96. The OA concentration contained in the sample was then calculated using the following equation:

$$x = \text{EXP} (y - b)/a$$

where x is the OA concentration in the sample (C_s) and y the absorbance of the sample.

The OA-toxin concentration in shellfish tissue was calculated as follows:

$$C_t (\mu\text{g}/\text{kg}) = (C_s (\text{nM}) \times \text{FD} \times \text{MW} (\text{g}/\text{mol}) \times V_e (\text{L}))/M_t (\text{g})$$

where C_t is the toxin concentration in tissue, expressed as equivalents of OA, FD is the methanolic extract dilution factor (31.25), MW is the OA molecular weight = 805, V_e is the methanolic extract volume (0.025 L), M_t is the tissue weight (5 g).

Samples with an OA concentration falling outside the working range (<0.5 nM or >2.8 nM) will be reported as <63 $\mu\text{g}/\text{kg}$ (or <0.5 nM) or >352 $\mu\text{g}/\text{kg}$ (or >2.8 nM), respectively.

2.5. Ruggedness Testing

The ruggedness testing was performed by introducing changes in the procedure and determining the effects on the sample quantification [14]. The variations used were chosen according to the values expected under normal laboratory conditions.

2.6. Spiking Procedure

Samples were spiked with OA Certified Reference Calibration Solution (NRC CRM-OA-c). The reference solution was prediluted to 2 μM in sample buffer and added accordingly. No Certified Reference Materials were available for DTX-1 and DTX-2 at the time of the performance testing. These toxins were first dissolved in methanol and diluted to 2 μM in sample buffer before adding to the samples.

A Certified Reference Material (NRC CRM-DSP-MUS-b) was also tested. However, the certified concentration of this material is far above the working range of the assay and the sample had to be diluted with blank mussel or king scallop. To do this, an amount of reference material was added as precisely as possible to 50 mL tubes, and weighed. The blank material was added on top and the mixture weighed again. Then, the amount of the mussel reference material per sample was calculated. This value was used as the theoretical spiked amount. The samples were analyzed with and without hydrolysis, as the reference material was only certified for OA and DTX-1, but ester derivatives of the OA-toxins could also be present as indicated in the CRM certificate. The total recovery was calculated according to the AOAC Official methods of analysis [15].

2.7. Method Comparison

A method comparison was also carried out with OkaTest, the mouse bioassay (MBA) and LC-MS/MS, using EU harmonized protocols for the last two methods [16,17].

Shellfish samples were previously tested by a third party laboratory using mouse bioassay (MBA) and LC-MS/MS, and kindly donated to do the method comparison.

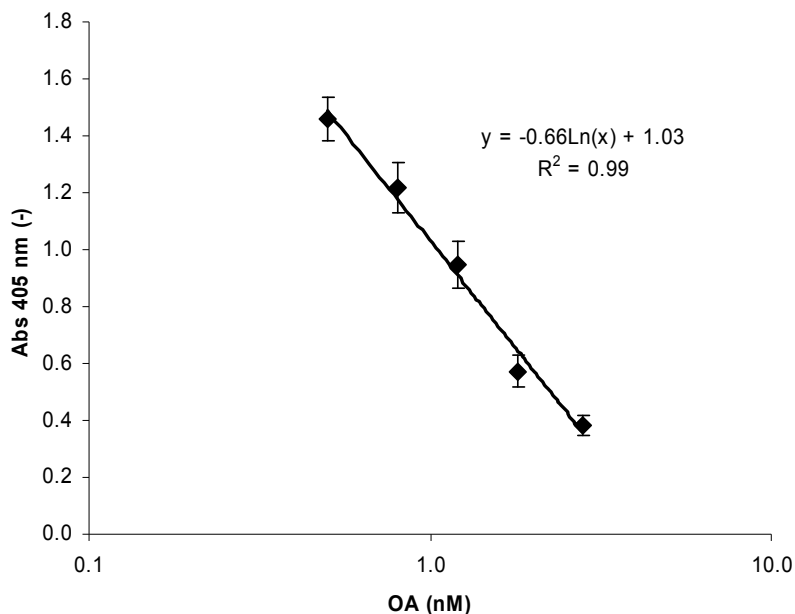
As MBA is a qualitative method, results obtained by OkaTest and LC-MS/MS were interpreted qualitatively for comparison purposes. Therefore, samples with a concentration ≥ 160 $\mu\text{g}/\text{kg}$ were regarded as positive, while samples with a concentration < 160 $\mu\text{g}/\text{kg}$ were reported negative.

3. Results and Discussion

3.1. Calibration of the Assay

The assay is calibrated by five OA standards prepared by dilution from the NRC CRM-OA-c with a concentration between 0.5 and 2.8 nM OA. Following the kits sample preparation (see material and methods), this will result in a working range between 63 and 352 $\mu\text{g}/\text{kg}$. Figure 1 shows a typical calibration curve from 5 different assays using different phosphatase batches. All calibration curves were evaluated according to the Pearson correlation coefficient obtained after a logarithmic fitting procedure ($r^2 > 0.96$).

Figure 1. Typical calibration curve of OkaTest produced as the mean of 5 phosphatase batches. The Pearson correlation coefficient (r^2) of the logarithmic fit was >0.96 for each batch. The figure shows the equation and r^2 of the mean. The error bars were calculated as ± 1 SD.



The bias introduced by the logarithmic fitting procedure on the calibration curve of the kit was estimated by recalculating the concentration of the OA dilutions using its own standard curve. The relative absolute difference was then calculated as the absolute difference between the theoretical and calculated OA concentration divided by the theoretical OA concentration and multiplied by 100 (Table 1). The best accuracy was found at levels around the regulatory limit (0.8% at 1.2 nM OA standards equals 151 μg OA equivalents/kg mollusk), while below that level (0.5 nM of OA), a 9.0% overestimation was calculated. Only minor deviations were calculated over the legal limit.

Table 1. Bias introduced due to the fitting procedure. Relative absolute difference was calculated from mean of 5 standard curves by relating the absolute difference to the theoretical OA concentration.

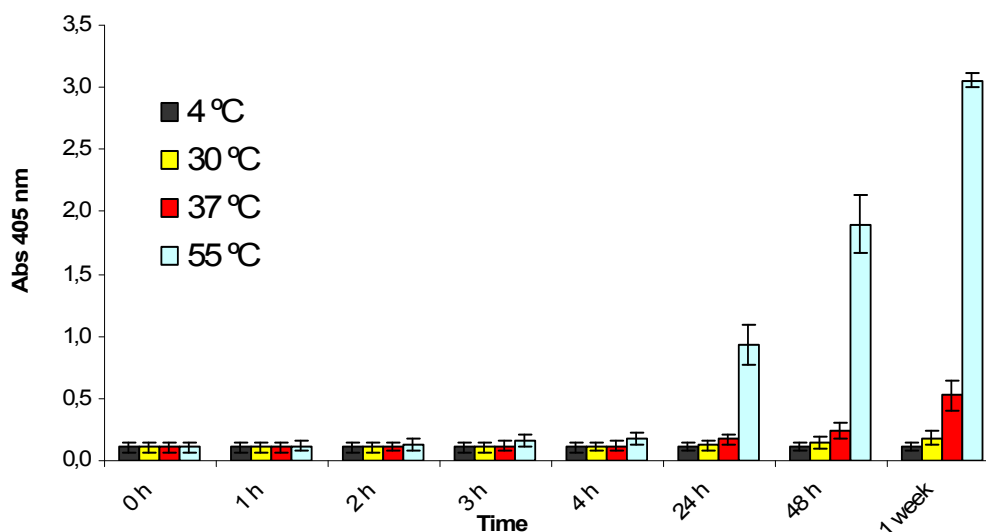
OA theoretical (nM)	OA calculated (nM)	Relative Absolute Difference
0.50	0.55	9.0%
0.80	0.83	3.8%
1.20	1.21	0.8%
1.80	1.78	1.1%
2.80	2.73	2.5%

3.2. Stability and Homogeneity of the Components

The stability and homogeneity of the critical components of the kit were studied by combining a real time and accelerated study design. Water soluble buffers such as the phosphatase dilution solution and the sample buffer were considered less critical, as sufficient internal know-how was available for these components and no stability problems were expected. Other components, such as the

ready-to-use chromogenic substrate, the PP2A or the OA standards, were specially developed for the phosphatase inhibition assay and were more extensively tested. Reagents were normally analyzed within the assay system or by performing specific tests depending on their particular characteristics. The ready-to-use substrate performed correctly in the OkaTest assay when stored for a year at temperatures between 2 and 15 °C (results not shown), as the background absorbance remained acceptable (below 0.3 absorbance units). However, accelerated studies showed that the substrate is sensitive to higher temperatures (Figure 2). After 24 h at 55 °C, the substrate was strongly hydrolyzed and after 1 week at 37 °C the absorbance of the substrate was above 0.6. Nevertheless, these results indicate that although the hydrolysis rate increases with temperature, it is very stable at temperatures below 15 °C and no problems should be expected under normal conditions of usage and storage.

Figure 2. Study of the temperature stability for the ready-to-use chromogenic substrate (p-Nitrophenyl phosphate). Absorbance at 405 nm was measured at different times and temperatures. Assays were performed in triplicate. The error bars were calculated as ± 1 SD.



The OA standards and the PP2A were estimated the most critical components, as their quantity and quality establish the working range and, to a great extent, the ruggedness of the assay. The enzyme quantity determines the amount of analyte that is needed for inhibition, while the enzyme quality assures the amount of product formed per time unit [18]. Likewise, the lack of stability or impurities of the OA standards directly affect the quantification, either overestimating, in the case of degradation of the OA, or underestimating, when impurities that can inhibit the PP2A are present. Therefore, greater emphasis was put on these components and the ‘between batch homogeneity’ was evaluated besides the stability of the components. The between batch homogeneity was studied by taking 1 set of standards or 1 vial of phosphatase from 5 different batches. These batches were chosen along the estimated shelf life of the compounds and tested in one single assay together with internal control samples. All batches performed according to the assays’ specifications ($r^2 > 0.96$) and the relative standard deviation was far below 15%, the expected value for samples assayed under repeatability conditions [19]. These results proved the stability of the enzyme for over 12 months at 4 °C and the homogeneity of between all batches tested (Table 2).

Table 2. Phosphatase stability and homogeneity. Five different phosphatase batches were tested at different stages of shelf life. Mean, standard deviation (SD) and relative standard deviation (RSDr) were calculated. Three internal control samples were used to verify correct quantification.

PP2A batch (shelf life)	Sample 1 (µg/kg)	Sample 2 (µg/kg)	Sample 3 (µg/kg)
1 (2 months)	95	160	310
2 (4 months)	100	169	304
3 (8 months)	88	162	323
4 (10 months)	94	156	300
5 (12 months)	90	144	341
mean	93	158	316
SD	5	9	17
RSDR	4.8%	6.0%	5.2%

For the OA standards, the same strategy was used. Five batches, covering 90% of the shelf life of the component (6 months), were tested in one assay to be able to single out the variation due to the standards' stability and homogeneity (Table 3). A sample shown to be blank (0 nM) was included to be able to calculate the effect of variables other than OA. The RSDr calculated from the absorbance values were all <3%, proving the stability and homogeneity of the standards over 6 months.

Table 3. OA standards stability and homogeneity. Five different batches of OA standards were tested at different stages of shelf life. The absorbances (405 nm) obtained for each of the standards are shown. Mean, standard deviation (SD) and relative standard deviation (RSDr) of these absorbances were calculated.

Standards OA (nM)	Absorbance 405 nm					mean	SD	RSDr
	batch 1 5 months	batch 2 4 months	batch 3 3 months	batch 4 2 months	batch 5 1 week			
0.0	2.042	2.100	2.064	2.073	2.120	2.079	0.031	1.5%
0.5	1.622	1.614	1.649	1.625	1.678	1.637	0.026	1.6%
0.8	1.462	1.390	1.386	1.375	1.372	1.397	0.037	2.7%
1.2	1.124	1.116	1.101	1.092	1.134	1.113	0.017	1.5%
1.8	0.772	0.792	0.769	0.822	0.809	0.793	0.023	2.9%
2.8	0.619	0.646	0.606	0.637	0.613	0.624	0.017	2.7%

3.3. Ruggedness

Enzymatic assays, such as OkaTest, can be sensitive to environmental factors, such as temperature, incubation time or reagent volume. To determine the impact of these factors, samples with concentrations around the regulatory limit were quantified at normal and suboptimal conditions (Table 4). The effect of temperature was tested by performing the OkaTest assay at three different temperatures 28, 30 and 32 °C, obtaining a RSD of 1.0%. These results showed that temperature variations of 2 °C did not affect the performance as RSDr values were lower than 10% usually obtained in the assay (Table 5).

Duration and pipetting volumes were evaluated alike and none of the variables affected the results of the test, with the exception of large pipetting errors. Pipetting errors of 5 μL in samples or phosphatase addition (errors of 10% and 7.1%, respectively) gave RSDr values of 14% and 17%, respectively. Precision in substrate addition was less critical. Pipetting samples and phosphatase are, however, the main sources of variability affecting PPIA and therefore care should be taken when adding these components.

Table 4. Ruggedness testing. The effects of variations of the normal assay conditions on sample quantification are shown.

Variable	Normal value	Variation	Mean value ($\mu\text{g}/\text{kg}$)	RSDr
Temperature	30 $^{\circ}\text{C}$	± 2 $^{\circ}\text{C}$	175	1.0%
Pre-incubation	20 min	18, 20, 22, 24 min	158	3.6%
Incubation	30 min	27, 30, 33, 36 min	147	2.9%
Syst. pipetting error	50, 70, 90 μL	± 2 μL	155	4.3%
Random pipetting error				
Sample	50 μL	± 5 μL	151	14%
PP2A	70 μL	± 5 μL	153	17%
Substrate	90 μL	± 5 μL	158	6.1%
Phosphatase solubility time	60 \pm 5 min	± 30 min	158	5.0%

Table 5. Intermediate precision of ten different mussel and scallops samples. Mean, standard deviation (SD), relative standard deviation (RSDr) were calculated. < 63: below the working range of the assay (63–352 $\mu\text{g}/\text{kg}$).

Sample	Origin	Day 1 ($\mu\text{g}/\text{kg}$)	Day 2 ($\mu\text{g}/\text{kg}$)	Day 3 ($\mu\text{g}/\text{kg}$)	Mean	SD	RSDr
1	Mussel	211	227	187	208	20	9.5%
2	Mussel	122	132	113	122	10	7.8%
3	Scallop	<63	<63	<63	-	-	-
4	Mussel	82	94	90	88	6	7.0%
5	Mussel	196	196	215	202	11	5.2%
6	Scallop	<63	<63	<63	-	-	-
7	Mussel	<63	<63	<63	-	-	-
8	Scallop	125	108	117	117	8	7.0%
9	Mussel	250	253	281	261	17	6.5%
10	Mussel	277	279	289	282	7	2.4%

3.4. Applicability

There are numerous descriptions of the application of protein phosphatase inhibition assays for determination of OA and its derivatives [7–13]. However, the inhibition pattern of OA, DTX1 and DTX2 is different and is supposed to correspond to their toxicity. One way to evaluate the inhibition capacity of toxins on an enzyme is by determining the IC_{50} , the concentration of toxin able to inhibit 50% of the maximum enzyme activity. This concentration depends, among others, on the amount of enzyme and the substrate concentration present in the assay [20] and therefore the IC_{50} values published for these toxins are difficult to compare [7,8,12,18,21,22]. The IC_{50} values found in our study were 1.2 nM for both OA and DTX-2, and 1.6 nM for DTX-1 (Figure 3) and are in accordance

with the ones obtained recently by Huhn *et al.*, 2009 [21]. However, these do not exactly correspond to the toxicity factors (TEF) that are used in analytical methods such as LC-MS/MS; as OA and DTX-1 have a TEF of 1, while DTX-2 has a TEF of 0.6, indicating equal toxicity for DTX-1 and OA and less toxicity for DTX-2 [2]. According to these values, our results would lead to an overestimation of the amount of DTX-2 and an underestimation of the amount of DTX-1 when compared with methods such as LC-MS/MS. However, the recovery data obtained for both DTX-1 and DTX-2 were similar to the ones obtained for OA (Table 6) suggesting that difference has a low impact in the determination of the level of toxins in shellfish samples.

Figure 3. Phosphatase inhibition curve obtained with okadaic acid (OA), dinophysistoxin-1 (DTX-1) and dinophysistoxin-2 (DTX-2). Each point is the mean obtained from three different phosphatase batches. The standard deviation is not shown to maintain the figure legible. The IC₅₀ values were 1.2 nM for both OA and DTX-2, and 1.6 nM for DTX-1.

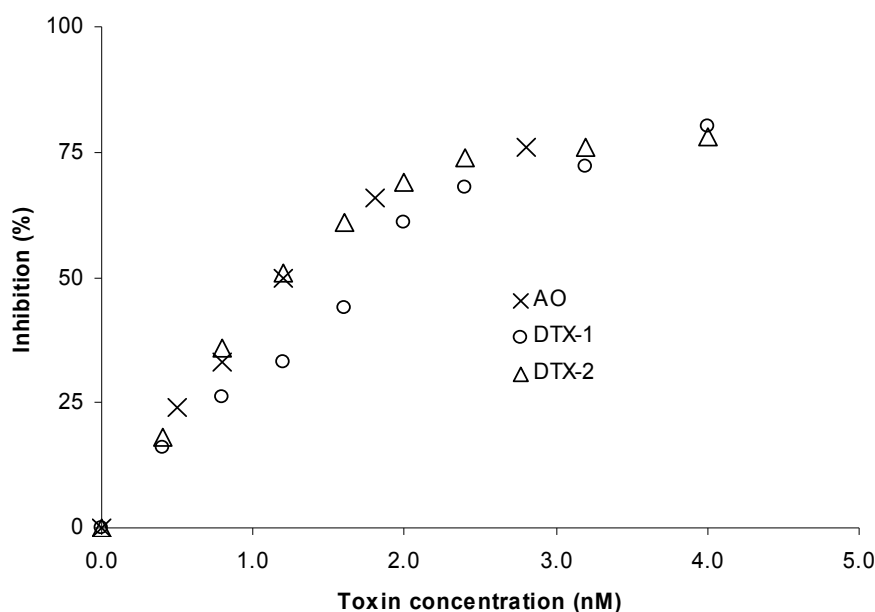


Table 6. Recovery of the different toxins was calculated testing 5 samples at 0.5, 1 and 1.5 times the regulatory limit on 3 different days. OA Certified Reference Material (NRC CRM-OA-c) was spiked on mussel and king scallop. DTX-1 and DTX-2 were spiked on king scallop. ND: not determined.

Toxin	Matrix	Recovery (RSDr)		
		80 µg/Kg	160 µg/Kg	240 µg/Kg
OA	Mussel	101% (15%)	90% (8.9%)	78% (5.4%)
	King scallop	114% (9.9%)	98% (8.4%)	106% (8.7%)
DTX-1	King scallop	102% (15%)	79% (12%)	88% (17%)
DTX-2	King scallop	93% (2.3%)	ND	ND

3.5. Limit of Detection, Limit of Quantification, Repeatability and Reproducibility

The limit of detection (LOD) and limit of quantification (LOQ) were determined by using a blank +3 SD or blank +10 SD approach [14]. For blank mussel material, the LOD and LOQ were 44 and

56 µg/kg, respectively. These values are both below the working range of the test and sufficiently below the current European legal limit of 160 µg/kg.

To estimate the precision, the assay was tested both under repeatability and intermediate precision conditions. The repeatability characteristics were estimated by analyzing 8 fractions of two naturally contaminated mussel samples and RSDr of 1.4% with a mean of 276 µg/kg, and 3.9% with a mean of 124 µg/kg were obtained (results not shown). The intermediate precision of the test was estimated by analyzing 7 samples with OA-toxin levels covering the working range of the assay on three different days by the same analyst. For all samples, the RSDr was well below the 15% RSDr limit as calculated by Horwitz [19]. Three samples tested as negative by LC-MS/MS were included to evaluate the consistency of the negative results (Table 5).

3.6. Accuracy

The accuracy of the method was estimated by calculating recoveries for OA, DTX-1 and DTX-2 and by testing a Certified Reference Material (NRC-CNRC). Five portions containing 5 grams of mussel or king scallop were spiked with one of the three toxins at 0.5, 1 and 1.5 times the regulatory limit (80, 160 and 240 µg/kg), except for DTX-2 that was only added up to a concentration of 80 µg/kg. The five portions were analysed on three different days to determine the intermediate precision characteristics of the test. OA recoveries between 78 and 101% in mussel and 98 and 114% in king scallop were obtained. RSDr values for this toxin were below or equal to 15%. Similar recoveries were obtained for the other two toxins (Table 6). These recoveries are in agreement with the 75 to 120% range that is expected for this concentration range [19]. The RSDr results in this study were higher than the ones obtained in the precision experiments (Table 4), specially for DTX-1. This might be a consequence of the spiking. As mentioned before, the higher IC₅₀ for DTX-1 compared to OA and DTX-2 had a low impact on the recovery.

Finally, four aliquotes of blank samples were spiked with the Certified Reference Material. The methanolic extract obtained was analysed with and without hydrolysis, and the recovery was estimated using the DTX-1 and OA content reported for the certified material. The recovery for the non-hydrolysed samples ranged from 71% to 98%, with a mean of 87% for mussle and 91% for king scallop (Table 7). These are acceptable recoveries and in accordance with the results showed in Table 6. However, the mean recovery of the hydrolysed samples was a 146% and 163% for mussle and king scallop, respectively. These percentages were far above the expected content of OA-toxins indicated in the reference material [23]. This could be due to the fact that the material is only certified for OA and DTX-1. Other esters of OA and DTX are reported in the certificate of anlysis for this material.

Table 7. Recovery experiment with Certified Reference Material (NRC CRM-DSP-MUS-b). Samples were analysed with and without hydrolysis.

Matrix	Spiked level (µg/kg) (n)	Without hydrolysis		With hydrolysis	
		Recovery	RSDr	Recovery	RSDr
mussel	219 (4)	87%	14%	146%	12%
king scallop	180 (4)	91%	5.0%	163%	2.8%

3.7. Method Comparison

A method comparison among MBA, LC-MS/MS and OkaTest was performed with a total of 37 samples. Results were compared qualitatively for all three methods and quantitatively between OkaTest and LC-MS/MS. The 160 µg/kg regulatory limit was used to decide whether the samples were positive or negative (Table 8).

Table 8. Methods comparison. Results from OkaTest, MBA and LC-MS/MS. 31 of the 37 samples were tested by MBA. Positive results (+): ≥ 160 µg/kg. Negative results (-): < 160 µg/kg. LOQ. Limit of quantification. NA: not available.

ID	M	MBA	LC-MS/MS	OKATEST	LC-MS/MS	OKATEST
1	Cockle	-	-	-	<LOQ	<LOQ
2	Cockle	+	+	+	193	252
3	Donax	-	-	-	82	97
4	Mussel	+	+	+	502	232
5	Mussel	+	-	+	<LOQ	268
6	Mussel	+	+	+	604	>352
7	Mussel	+	+	+	894	>352
8	Mussel	+	+	+	414	306
9	Mussel	+	+	+	444	>352
10	Mussel	NA	-	-	<LOQ	<LOQ
11	Mussel	NA	+	+	357	>352
12	Mussel	NA	-	-	<LOQ	<LOQ
13	Mussel	NA	-	-	<LOQ	<LOQ
14	Mussel	-	-	-	<LOQ	122
15	Mussel	+	-	+	158	196
16	Mussel	+	+	+	177	250
17	Mussel	+	+	+	288	265
18	Mussel	+	+	+	202	196
19	Mussel	+	+	+	390	277
20	Mussel	+	+	+	658	305
21	Mussel	+	+	+	392	310
22	Mussel	+	+	+	329	315
23	Mussel	+	+	+	232	270
24	Mussel	+	+	+	235	277
25	Mussel	+	-	-	152	135
26	Mussel	+	-	+	98	164
27	Mussel	+	+	+	168	211
28	Mussel	+	+	+	209	251
29	Mussel	+	-	+	113	191
30	Mussel	NA	+	-	292	<LOQ
31	Mussel	NA	+	+	316	304
32	Mussel	-	-	-	<LOQ	<LOQ
33	Mussel	+	+	-	177	124
34	Mussel	+	+	+	247	216
35	Mussel	+	+	-	185	144
36	Scallop	+	+	+	184	264
37	Scallop	-	-	-	<LOQ	<LOQ

In general, the qualitative interpretation of the results indicates that the three methods obtained equivalent results, especially taking into account that these are conceptually different methods. The OkaTest disagreed with both MBA and LC-MS/MS on two occasions (samples 33 and 35). OkaTest detected levels of OA-toxins in those two samples, but below the EU regulatory limit (124 and 144 µg/kg), while the samples were positive according to the other two methods). A third sample (25) was also identified as negative by OkaTest and positive by MBA. LC-MS/MS also gave a negative result for sample 25. The concentration of this sample determined by both methods was just below the EU regulatory limit.

The LC-MS/MS differed on four occasions: all four negative according to LC-MS/MS, but positive by the other two methods. Three of the samples (15, 26 and 29) contained OA-toxins below the EU regulatory limit, but sample 5 was quantified under the method's LOQ. Finally, one sample (30) was positive by LC-MS/MS, but under the LOQ by OkaTest. Sample 30 was not tested by MBA due to lack of material.

Quantitative results obtained by LC-MS/MS and Okatest showed some differences. About two thirds of the samples gave similar results ($\pm 25\%$) with both methods, but the rest of the samples did not show a clear tendency. There is no evident explanation for this and further investigation would be required.

4. Conclusions

A colorimetric phosphatase inhibition assay for determination of OA-toxins, OkaTest, was single laboratory validated according to international methods validation guidelines. The limit of quantification of the method is well below the EU regulatory limit and the method permitted the easy quantification of up to 43 samples within one hour, excluding sample preparation. The method is robust, with very good precision characteristics, adequate specificity and accuracy.

This colorimetric phosphatase inhibition assay could be used as a complementary assay to the reference method for determination of lipophilic toxins, once a collaborative study has been completed and it has been successfully tested under recognized proficiency tests. This assay could be applied for monitoring purposes when OA-toxins are identified to be responsible for a bloom.

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Note: Collaborative efforts between the EURLMB and ZEU-INMUNOTEC do not amount to an endorsement of the firm's products.

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FOOD CHEMICAL CONTAMINANTS**Quantitative Determination of the Okadaic Acid Toxins Group by a Colorimetric Phosphatase Inhibition Assay: Interlaboratory Study****HENRY SMIENK and ELENA DOMÍNGUEZ¹**

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An interlaboratory collaborative study to validate a colorimetric phosphatase inhibition assay for quantitative determination of the okadaic acid (OA) toxins group in molluscs, OkaTest, was conducted. Eight test materials, including mussels, scallops, clams, and cockles, were analyzed as blind duplicates. Blank samples and materials containing different OA toxin levels ranging from 98 to 275 µg/kg OA equivalents were included. The study was carried out by a total of 16 laboratories from 11 different countries. Values obtained for repeatability relative standard deviations (RSD_r) ranged from 5.4 to 11.2% (mean 7.5%). Reproducibility RSD (RSD_R) values were between 7.6 and 13.2% (mean 9.9%). The Horwitz ratio (HorRat) values ranged between 0.4 and 0.6. A recovery assay was also carried out using a sample spiked with OA. A mean recovery of 98.0% and an RSD of 14.5% were obtained. The results obtained in this validation study indicate that the colorimetric phosphatase inhibition assay, OkaTest, is suitable for quantitative determination of the OA toxins group. OkaTest could be used as a test that is complementary to the reference method for monitoring the OA toxins group.

O kadaic acid (OA) and its analogs dinophysistoxin-1 and -2 (DTX1, DTX2), together with their ester forms, are known as the OA toxins group. These lipophilic and heat stable toxins are produced by dinoflagellates and can be found in various species of shellfish, mainly in filter-feeding bivalve molluscs.

OA toxins causes diarrhetic shellfish poisoning, which is

characterized by symptoms, such as diarrhea, nausea, vomiting, and abdominal pain. These symptoms may occur in humans shortly after consumption of contaminated bivalve molluscs, such as mussels, clams, scallops, or oysters. Inhibition of serine/threonine phosphoprotein phosphatases (PPs) is assumed to be responsible for these toxic effects. These compounds are also involved in tumor promotion (1). Therefore, these toxins are regulated by European Union law.

Regulation (EC) No. 853/2004 (2) states that live bivalve molluscs placed on the market for human consumption must not contain marine biotoxins in total quantities (measured in the whole body or any part edible separately) that exceed 160 µg of OA equivalents/kg for OA, dinophysistoxins, and pectenotoxins together.

Commission Regulation (EC) No. 15/2011 (3) indicates that in the case of lipophilic toxins including OA toxins, LC/MS/MS is the reference method for routine testing of official controls or any checks done by food operators. This regulation has recently amended the Commission Regulation (EC) No. 2074/2005 (4), in which biological methods (mouse and rat bioassay) were considered the reference. From now on, they will only be used for a transitional period of time (until the end of 2014) or in special circumstances.

Both regulations (No. 2074/2005 and No. 15/2011) contemplate other methods for routine testing of lipophilic toxins, providing they are intralaboratory-validated and successfully tested under a recognized proficiency test scheme. Those methods should detect, either alone or in combination with others, all of the lipophilic toxin analogs (OA, pectenotoxins, yesotoxins, and azaspiracids group toxins). The protein phosphatase inhibition assay (PPIA) is specifically mentioned in these regulations as an alternative or complementary method, considering that the PPs are known to be OA-toxins natural targets (5, 6). In-house PPIAs using different phosphatase sources and colorimetric or fluorometric substrates have been previously developed (7–12). Later improvements to detect all OA derivatives by hydrolysis of samples were also suggested

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Table 1. Details of matrixes and species origin of test materials used in this study

Code	Matrix/Species	Origin
A	Mussel (<i>M. galloprovincialis</i>)	Galicia (NW Spain)
D	Clam (<i>V. pullastra</i>)	Food & Agricultural Organization, 37 Mediterranean Sea
E	Mussel (<i>M. galloprovincialis</i>)	Galicia (NW Spain)
F	Scallop (<i>P. maximus</i>)	FAO 27 NE Atlantic
G	Clam (<i>V. decussatus</i>)	Galicia (NW Spain)
K	Clam (<i>V. romboides</i>)	Galicia (NW Spain)
L	Cockle (<i>C. edulis</i>)	Portugal and Galicia (NW Spain)
N	Mussel (<i>M. edulis</i>)	Ireland
BM	Scallop (<i>P. maximus</i>)	Scotland

(13), and a collaborative study was also performed with a fluorometric PPIA (14). However, none of those assays was commercially available for routine analysis, nor were they demonstrated to comply with the legislation requirements.

ZEU-INMUNOTEC (Zaragoza, Spain) has developed a commercial kit (OkaTest, formerly Toxiline-DSP) based on a colorimetric PP2A inhibition assay for quantification of the OA toxins group in molluscs (15).

The PPIA described in this study uses a human PP2A purified by ZEU-INMUNOTEC that has showed higher sensitivity than other commercial and genetic engineering produced enzymes (16). PP2A was stabilized by freeze-drying to obtain a standardized assay with shelf life of up to 12 months at 4°C (15). Colorimetric substrate was chosen over a fluorometric one as the latter is less stable and, therefore, less appropriate for ready-to-use kits. Besides, fluorometric assays require specific equipment not often available in routine testing laboratories; therefore, they are difficult to use for monitoring purposes.

The robustness and performance of OkaTest were evaluated by the manufacturer in a single-laboratory validation according to AOAC and Eurachem guidelines (15). All of the results obtained showed that the OkaTest kit is robust and accurate, and, therefore, suitable for an interlaboratory study.

Interlaboratory Study

A colorimetric PPIA, OkaTest, was interlaboratory-validated for quantification of the OA toxins group. The main purpose of this study was to determine repeatability and between-laboratory reproducibility. A recovery assay was also carried out, and accuracy of the method confirmed.

A validation management team (David Clarke, Elena Domínguez, Katrin Kapp, Panagiota Katikou, and María Luisa Rodríguez) was appointed to supervise, advise on the accomplishment of the study, and ensure its independence. A total of 16 laboratories from 11 different countries in Europe and South America participated in the study.

The study plan including details of the test method, experimental design, preparation of test materials, instructions for participants, key personnel, schedule, and data analysis was prepared and agreed to by the validation management team.

Participants were fully informed of the study design prior to distribution of testing materials.

Eight different test materials, as blind duplicates, were analyzed by each laboratory on 2 different days. Five materials contained different OA toxin levels, all naturally contaminated except for one that was partially spiked. Three of the test materials were blank samples. An additional blank material (BM) was used in the recovery study. The test materials comprised four different genera of molluscs (*Mytilus spp.*, *Pecten spp.*, *Venerupis spp.*, and *Cerastoderma spp.*) and seven different species. Details of the materials used are shown in Table 1. The materials were prepared by the Spanish Association of Seafood Products Manufacturers (ANFACO-CECOPECA; Vigo, Spain) as explained below.

All participants sent back an electronic copy of a tailor-made Excel reporting sheet for each day of analysis with raw data and final results for each test material. The reporting sheets were checked upon receipt for obvious errors in sample codes and calculations.

Participants also completed a questionnaire with details of the equipment used and preparation of reagents and samples, as well as feedback on the assay.

Preparation of Test Materials

Materials A and E (mussel) and D and K (clam) were purchased from the retail market fresh and alive. They were thoroughly cleaned outside and inside with fresh water to remove sand and any other foreign materials. Tissues were removed from the shell, transferred to strainers, and drained for 5 min before homogenization (blender and Ultraturrax[®]; IKA, Staufen, Germany). The homogenate (at least 450 g) was then distributed into plastic containers (5.0 ± 0.1 g), frozen, and stored at -20 ± 2°C until analysis or the day of shipment.

Materials F (scallop) and G (clam) were purchased frozen

Table 2. Total concentration of OA toxins group (µg/kg) determined by OkaTest, and toxins profile by LC/MS/MS

Test material ^a	Matrix/species	Total OA equivalents, µg/kg ^b	OA toxins content ^c
BM	Scallop (<i>P. maximus</i>)	<LOD	—
A	Mussel (<i>M. galloprovincialis</i>)	<LOD	—
F	Scallop (<i>P. maximus</i>)	<LOD	—
G	Clam (<i>V. decussatus</i>)	<LOD	—
E	Mussel (<i>M. galloprovincialis</i>)	79 ± 5	OA
L	Cockle ^d (<i>C. edulis</i>)	168 ± 11	OA, DTX1, and DTX2
D	Clam (<i>V. pullastra</i>)	240 ± 9	OA
K	Clam (<i>V. romboides</i>)	250 ± 6	OA
N	Mussel ^e (<i>M. edulis</i>)	276 ± 6	OA and DTX2

^a Samples presented in increasing order of concentration.

^b Determined by OkaTest; LOD = 44 OA equivalents µg/kg.

^c Determined by LC/MS/MS.

^d Artificially contaminated with DTX1 and mixed with blank material.

^e Mixed with blank material.

Table 3. Results from homogeneity study for test materials for the determination of OA (µg OA total equivalents/kg)

Test material	Variance of sums, V_s	Analytical variance, s_{an}^2	Allowable sampling variance, σ_{all}^2	Sampling variance, S_{sam}^2	Critical value, c	Test for homogeneity result
D	166	90.7	36.8	116	310	$S_{sam}^2 < c$
E	84.7	8.09	19.8	11.1	29.1	$S_{sam}^2 < c$
K	139	19.6	32.5	126	257	$S_{sam}^2 < c$
L	356	46.9	85.7	55.6	152	$S_{sam}^2 < c$
N	124	24.2	28.4	154	314	$S_{sam}^2 < c$

from the retail market. They were thawed at room temperature, cleaned, and prepared as described above.

Material L (cockle) was provided cleaned, blended, and frozen by the European Reference Laboratory for Marine Biotoxins (EURLMB, Vigo, Spain). The sample contained OA, DTX2, and traces of DTX1. In order to achieve a suitable toxin profile, the sample was mixed with fresh cockle from the same species (*C. edulis*) without toxin prior to being spiked with DTX1 (Wako Chemicals, Neuss, Germany). The sample was thawed at room temperature, mixed with the cockle blank material (purchased in Porto, Portugal), and spiked. Then, it was distributed into plastic containers (5.0 ± 0.1 g), frozen, and stored at $-20 \pm 2^\circ\text{C}$ until the day of shipment.

Material N (mussel) was provided cleaned, blended, and frozen by the National Reference Laboratory of Ireland, Galway, Ireland. The sample contained a high level of OA toxins, so it was mixed with mussel (*M. edulis*) without toxin (purchased in a retail market in Ireland) to achieve a suitable toxin concentration. The sample was thawed at room temperature, mixed, and distributed into plastic containers (5.0 ± 0.1 g). The material was then frozen and stored at $-20 \pm 2^\circ\text{C}$ until the day of shipment.

The BM (scallop) was provided blended and homogenized by Integrin Advanced Bioscience (Oban, Scotland) and stored frozen at approximately $-20 \pm 2^\circ\text{C}$ until the day of shipment.

Homogeneity and stability of test materials were studied according to the International Harmonized Protocol for the Proficiency Testing of Analytical Chemistry Laboratories (17). Ten containers of 5 g were randomly selected for each material. The content of each container was homogenized and extracted, and two test portions (from the sample extract) were analyzed to estimate the analytical variance. A total of 20 portions/material

were tested under repeatability conditions and in a random order using the OkaTest kit.

To ensure the stability of the materials during shipment to participants and the study duration, aliquots of each material were taken randomly and split into two subsets, each of them containing five samples. One subset was used as control and stored at $-18 \pm 1^\circ\text{C}$. The second was stored under experimental conditions of $9.0 \pm 1^\circ\text{C}$ for 5 days. Samples of both subsets were randomized before testing and analysis simultaneously using the OkaTest kit under repeatability conditions. The test materials were also analyzed by LC/MS/MS (18, 19) to determine the OA toxin profile.

The test materials were blind coded by EURLMB and distributed by ANFACO-CECOPESCA to the participants. The codes were securely kept by EURLMB until statistical analysis was carried out.

The materials were shipped in isothermal boxes with dry ice and were received within the following 2 days by most participants. Materials sent to South American countries were delivered more than a week after the dispatch date, as they have long customs check up procedures. Samples were, however, reported to have been kept frozen while stored at customs. Two laboratories informed that the box containing the samples did not arrive in good conditions, and six reported that samples were cold, but defrosted.

PPIA

Principle

OkaTest is an enzymatic test based on a colorimetric PPIA for quantitative determination of OA and other toxins of the OA group, including DTX1, DTX2, and their ester forms.

Table 4. Results obtained for the stability assays conducted for materials D, E, K, L, and N

Test material	Storage conditions		Absolute difference D	Variance <i>F</i> -test	<i>t</i> -test	Test criterion	
	-18 ± 1°C	9.0 ± 1°C				C	D < C
	Mean						
Total OA equivalents, µg/kg							
D	265 ± 10	262 ± 15	3.02	0.54	0.71	34.5	Pass
E	84.0 ± 4	85.1 ± 3	-1.19	0.45	0.62	10.9	Pass
K	255 ± 8	257 ± 7	-1.57	0.87	0.75	33.2	Pass
L	171 ± 7	169 ± 8	1.63	0.79	0.73	22.2	Pass
N	343 ± 24	355 ± 32	-13.0	0.58	0.49	44.6	Pass

Table 5. Calibration curve parameters obtained by each laboratory every day of the study

Lab	R ²		Slope		Absorbance 405 nm, lowest standard 0.5 nM		Absorbance 405 nm, highest standard 2.8 nm	
	Day 1	Day 2	Day 1	Day 2	Day 1	Day 2	Day 1	Day 2
A	0.99	0.98	-0.12	-0.45	0.734	1.287	0.524	0.505
B	0.99	0.99	-0.50	-0.65	1.157	1.425	0.334	0.339
C	0.98	0.98	-0.64	-0.44	1.530	1.177	0.496	0.468
D	0.98	0.98	-0.67	-0.58	1.537	1.402	0.430	0.459
E	0.97	0.98	-0.51	-0.48	1.222	1.221	0.409	0.436
F	1.00	0.99	-0.72	-0.74	1.684	1.726	0.482	0.491
G	0.98	1.00	-0.79	-0.58	1.781	1.411	0.462	0.423
H	0.99	0.99	-0.78	-0.73	1.644	1.609	0.366	0.414
I	0.99	0.99	-0.76	-0.68	1.661	1.486	0.409	0.357
J	0.97	0.98	-0.41	-0.45	1.164	1.204	0.498	0.458
K	0.99	0.98	-0.77	-0.74	1.712	1.690	0.438	0.485
L ^a	0.93	0.96	-0.63	-1.13	1.488	2.588	0.425	0.709
M	0.99	0.99	-0.78	-0.65	1.697	1.464	0.419	0.390
N	0.99	0.98	-0.54	-0.65	1.273	1.497	0.384	0.444
O	0.97	0.98	-0.49	-0.32	1.188	0.992	0.396	0.470
P	0.97	0.99	-0.27	-0.58	1.015	1.474	0.549	0.520

^a Standard curve obtained by Laboratory L on Day 1 was rejected as R² criterion was not met. Assay could not be repeated due to time issues.

This method is applicable to shellfish species, such as mussels, clams, cockles, and scallops.

The toxicity of the OA toxins group is directly related to its inhibitory activity against a family of structurally related PPs, in particular PP1 and PP2A. OkaTest uses this strong inhibitory activity to determine the OA content in shellfish using the PP2A with a chromogenic substrate for this enzyme. After the substrate's hydrolysis by the enzyme, the product can be measured at 405 nm by a microplate reader. As the ability of the PPs to hydrolyze the substrate depends on the amount of OA and analogs in the samples, the toxin concentration can be calculated by using a standard curve.

Apparatus

(a) *Micropipets*.—Adjustable 100, 200, and 1000 μ L (Thermo LabSystems, Helsinki, Finland).

(b) *Ultra homogenizer*.

(c) *Block heater or incubator*.—For $30 \pm 2^\circ\text{C}$ (ZEU-INMUNOTEC, Zaragoza, Spain).

(d) *Microwell absorbance reader*.—405 \pm 10 nm wavelength filter (Thermo LabSystems).

(e) *Water bath*.—Set at $76 \pm 2^\circ\text{C}$ (Raypa, Barcelona, Spain).

(f) *Centrifuge tubes*.—Graduated 50 mL.

(g) *Laboratory glassware*.

Reagents

(a) *Extraction solvent*.—Methanol, reagent grade, 100% (v/v; Sharlab, Barcelona, Spain).

(b) *HCl*.—Reagent grade, 37% (v/v; Sharlab).

(c) *NaOH*.—Reagent grade (Sharlab).

(d) *Deionized water*.—Type II, ISO 3696 (Ellix 5; Millipore, Germany).

(e) *OkaTest kit*.—From ZEU-INMUNOTEC containing:

(1) 96-well microtiter plate and plate adhesive film.

(2) Lyophilized PP2A purified from human blood cells.

(3) Ready-to-use OA Standards of 0.5, 0.8, 1.2, 1.8, and 2.8 nM, prepared from the OA reference solution (NRC CRM-OA-c, Institute for Marine Biosciences, Halifax, Canada).

(4) Chromogenic substrate.

(5) Phosphatase dilution buffer.

(6) Stock buffer solution.

(7) OA Spiking solution (2 μ M) prepared from the OA reference solution (NRC CRM-OA-c, Institute for Marine Biosciences).

Spiking Procedure

Due to the limited experience on the homogeneity and stability of spiked samples with OA toxins, each participant prepared a spiked sample on the day of the assay. A BM and an OA solution of known concentration (2 μ M, to prepare a final concentration of 161 μ g/kg) were provided to each participant.

A blank sample was spiked with OA solution for the recovery study as follows:

(a) Mix 500 μ L OA spiking solution (2 μ M) with 5.0 ± 0.1 g homogenous blank sample.

(b) Add 25 mL extraction solvent [methanol, 100% (v/v)] to the mixture and shake for 2 min by vortexing. Proceed with the extraction procedure described below under point (b).

Sample Extraction

(a) Thaw each aliquot with 5.0 ± 0.1 g homogenized mollusc at room temperature ($22 \pm 2^\circ\text{C}$). Add 25 mL extraction solvent [methanol, 100% (v/v)]; then mix for 2 min using an ultra homogenizer.

(b) Centrifuge at 2000 g for 10 min at 4°C. The supernatant is called “methanolic extract.”

(c) Pipet 640 µL methanolic extract into a 50 mL graduated centrifuge tube and add 100 µL 2.5 M NaOH.

(d) Seal the test tube and heat at 76 ± 2°C for 40 min in a water bath.

(e) Do not cool the sample; add 80 µL 2.5 M HCl immediately.

(f) Add 19.18 mL buffer solution with a glass pipet up to a total volume of 20 mL.

Assay Procedure

(a) Rehydrate the lyophilized phosphatase (PP2A) by adding 2.0 mL phosphatase dilution buffer to the vial and mix gently for 60 ± 5 min at room temperature (22 ± 2°C) on a roller mixer or a shaker (maximum 60 rpm) (both from JP Selecta, Barcelona, Spain).

(b) Add 50 µL each sample extract or standard to wells. Samples and standards have to be analyzed in duplicate.

(c) Add 70 µL phosphatase solution to each well. Cover the plate with the adhesive film provided in the kit, and mix by gentle tapping on the side.

(d) Incubate at 30 ± 2°C for 20 ± 0.5 min.

(e) Remove the adhesive film and add 90 µL chromogenic substrate to each well and mix by tapping gently on the side. Incubate at 30 ± 2°C for 30 ± 0.5 min.

(f) Read the absorbance of samples and standards at 405 ± 10 nm.

Calculations

The results were calculated from a standard curve by plotting the absorbance values on a linear y axis and the concentration of OA on a logarithmic x axis, and using a logarithmic fitting. As an acceptability criterion for the assay, the Pearson correlation coefficient R² had to be equal to or greater than 0.96. The OA concentration contained in the sample was then calculated using the following equation:

$$x = \text{EXP} (y - b)/a$$

where *x* is the OA concentration in the sample (*C_s*), *y* the absorbance of the sample, *a* is the slope, and *b* is the y-intercept.

The OA toxin concentration in shellfish tissue was calculated as follows:

$$C_t, \mu\text{g}/\text{kg} = [C_s (\text{nM}) \times \text{FD} \times \text{MW} (\text{g}/\text{mol}) \times V_e (\text{L})] / M_t (\text{g})$$

where *C_t* is the toxin concentration in tissue expressed as equivalents of OA, *FD* is the methanolic extract dilution factor, *MW* of OA = 805, *V_e* is the methanolic extract volume (0.025 L), and *M_t* is the tissue weight (5 g).

Samples with an OA concentration falling outside the working range (<0.5 nM or >2.8 nM) will be reported as <63 µg/kg (or <0.5 nM) or >352 µg/kg (or >2.8 nM), respectively.

Results were recorded by each participant in a tailor-made Excel spreadsheet with which the results were automatically calculated when the absorbance values were entered. All participants sent back an electronic copy of the reporting sheet for each day of analysis.

Table 6. Individual results (µg OA total equivalents/kg) reported from laboratories A to P for Materials A, D, E, F, G, K, L, and N on Days 1 and 2. Invalid or incorrect results are those in bold type.

Lab	µg OA total equivalents/kg															
	Material															
	A		D		E		F		G		K		L		N	
	Day															
	1	2	1	2	1	2	1	2	1	2	1	2	1	2	1	2
A	<63	<63	186	239	97	102	<63	<63	<63	<63	248	281	167	174	210	247
B	<63	<63	251	266	100	101	<63	<63	<63	<63	302	299	177	190	273	277
C	<63	<63	244	233	96	87	<63	<63	<63	<63	279	246	174	160	256	251
D	<63	<63	264	253	125	100	<63	<63	<63	<63	282	277	189	223	269	295
E	<63	<63	210	233	101	120	<63	<63	<63	<63	239	244	156	181	226	219
F	<63	<63	252	250	113	116	<63	<63	<63	<63	287	286	166	165	271	275
G	<63	<63	246	252	89	100	<63	<63	<63	<63	356 ^a	269 ^a	192	192	274	236
H	<63	<63	253	250	90	99	<63	<63	<63	<63	291	301	175	179	271	270
I	<63	<63	252	254	95	87	<63	<63	<63	<63	284	283	169	161	265	253
J	70 ^a	98 ^a	238	239	163 ^a	102 ^a	<63	<63	78 ^a	67 ^a	248	268	239	184	246	235
K	<63	<63	253	264	81	81	<63	<63	<63	<63	295	300	152	160	247	266
L	—	<63	—	242	—	145	—	<63	—	—	—	266	—	202	—	182
M	<63	<63	257	255	101	104	<63	<63	<63	<63	292	274	177	176	271	272
N	<63	<63	261	251	98	101	<63	<63	<63	<63	285	285	161	181	257	250
O	<63	<63	221	223	91	94	<63	<63	<63	<63	270	249	179	184	259	244
P	<63	<63	192	241	69 ^a	153 ^a	<63	<63	<63	<63	226	278	97	173	206	259

^a Outlier.

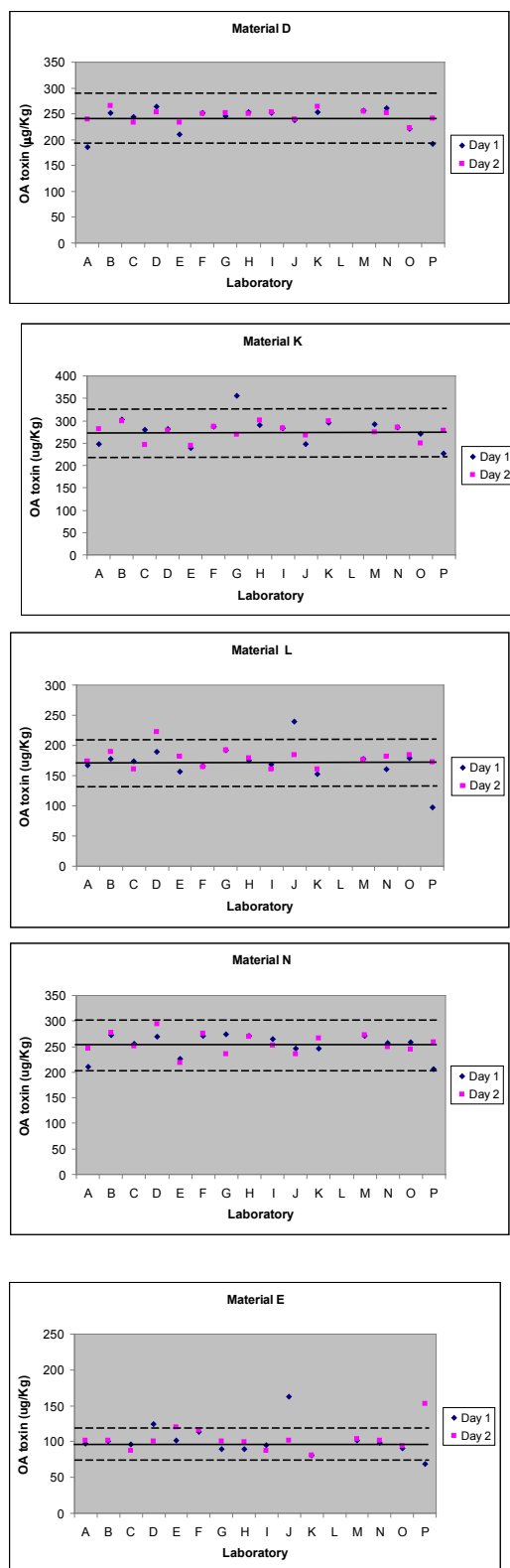


Figure 1. Individual results for each test material obtained per lab and per day of analysis (including outliers). The solid line shows the assigned mean value calculated in this study for each material. The dashed lines indicate the theoretical reproducibility SD determined for each material in this study (PRSD_R).

Statistics

Analysis of Valid Data and Outliers

Statistical data analysis was carried out following the approach described in the AOAC/IUPAC guidelines (17, 20). Submitted results were initially reviewed to remove invalid data. Results from assays with calibration curves with a $R^2 < 0.96$ and results outside the working range or showing deviations from the Standard Operating Procedure were considered invalid.

The valid data were first analyzed for possible outliers applying the Cochran and Grubbs tests. Then, precision parameters, HorRat values, and recovery were calculated.

The Cochran test was applied to remove laboratories showing significantly greater variability among replicate (within-laboratory) analyses than the other laboratories for a given material. A 1-tail test at a probability value of 2.5% was applied (17, 20).

The Grubbs test was used to remove results from laboratories with extreme averages (17, 20). This test was applied to the remaining values from the Cochran test. A single value test (two-tail, $P = 2.5\%$) was first applied, followed by a pair value test (two values at the highest end, two at the lowest end, and one at each end, at an overall $P = 2.5\%$).

Precision

To estimate the precision of the method, the within-laboratory repeatability and between-laboratory reproducibility were determined by calculating s_r (repeatability SD), s_R (reproducibility SD), RSDs (RSD_r and RSD_R), repeatability and reproducibility limits (r and R), and HorRat values. These parameters were calculated following the AOAC guidelines (20).

Recovery

For recovery calculations, the marginal recovery was calculated as follows:

$$\text{Recovery, \%} = 100 (C_f - C_u) / C_A,$$

where C_f is the amount found for the spiked concentration, C_u is the amount present originally for the unspiked concentration, and C_A is the amount added.

Results and Discussion

Test Material Results

The test materials were first analyzed by OkaTest and LC/MS/MS to determine the content and profile of OA toxins. Results obtained by both methods for samples A, F, and G showed concentration for OA toxins below their LOD (44 and 40 µg/kg, respectively). The BM was tested by LC/MS/MS (19) at EURLMB, and no peaks were detected for this group of toxins (LOD for this method is 15 µg/kg). Therefore, materials A, F, G, and BM were considered blank; therefore, no homogeneity or stability studies were carried out.

Analyses by LC/MS/MS were used to identify the toxin profile and to ensure that all toxins belonging to the OA group were present in the materials. Table 2 shows concentration

Table 7. Details of the test materials, number of results submitted, and results after removing outliers, together with performance values of precision (repeatability and reproducibility) obtained for the colorimetric OkaTest^a

Test material	Matrix	Runs/lab	No. labs submitting results	No. labs after invalid/incorrect results	No. of labs after outliers ^b	Mean (µg total equivalent OA/kg) ^c	Repeatability ^c			Reproducibility ^c			
							S _r	r	RSD _r %	µg total equiv.OA/kg			
										S _R	R	RSD _R %	HorRat
A	Mussel <i>M. galloprovincialis</i>	2	16	14	—	<63	—	—	—	—	—	—	—
D	Clam <i>V. pullastra</i>	2	16	15	15 (0)	242	14.7	41.2	6.1	19.4	54.4	8.0	0.4
E	Mussel <i>M. galloprovincialis</i>	2	16	15	13 (2)	98.8 (102)	7.32 (20.8)	20.5 (58.4)	7.4 (20.5)	10.7 (19.6)	30.0 (54.8)	10.7 (19.2)	0.5 (0.8)
F	Scallop <i>P. maximus</i>	2	16	15	—	<63	—	—	—	—	—	—	—
G	Clam <i>V. decussatus</i>	2	16	14	—	<63	—	—	—	—	—	—	—
K	Clam <i>V. romboides</i>	2	16	15	14 (1)	275 (277)	14.9 (21.4)	41.8 (60.1)	5.4 (7.7)	21.0 (25.0)	58.7 (70.1)	7.6 (9.0)	0.4 (0.5)
L	Cockle <i>C. edulis</i>	2	16	15	15 (0)	175	19.6	55.0	11.2	23.2	64.9	13.2	0.6
N	Mussel <i>M. edulis</i>	2	16	15	15 (0)	255	15.6	43.7	6.1	20.7	58.1	8.1	0.4

^a S_r = Repeatability SD, S_R = reproducibility SD, RSD_r = repeatability RSD, RSD_R = reproducibility RSD, r = repeatability limit, R = reproducibility limit.

^b Number of laboratories remaining after removal of outliers (number of outliers).

^c Mean, repeatability, and reproducibility (values obtained including outliers).

in OA equivalents determined by OkaTest and toxins profile of the different materials used. All test materials were found to be stable for the duration of the study and with sufficient homogeneity (Tables 3 and 4).

Interlaboratory Study Results

All participants who received test materials reported results. The sample concentration was calculated by standard curves obtained by each laboratory every day of analysis. Fit parameters of each standard curve are shown Table 5. Although the slopes show differences depending on the laboratory and day, the calculated samples concentration was not affected. The data obtained by each laboratory per test material and day of analysis are shown in Table 6.

All individual values obtained per material, day and laboratory were also plotted. One graph per material is shown in Figure 1. The solid lines represent the assigned mean value obtained for each material in this study (Table 7). The area between the dashed lines demonstrates the range of deviation from the mean value based on the theoretical reproducibility SD (PRSD_R).

Two laboratories reported one of the assays with R² < 0.96; one (Laboratory A) repeated the analysis obtaining R² within the required criterion. Laboratory L, however, could not repeat the assay on time, and those results were considered invalid and removed for statistical analysis.

Materials A, F, and G were not statistically analyzed, as they were blank samples. However, Laboratory J reported values within the working range of the test for Materials A and G. These values are considered incorrect according to the AOAC

guidelines (20), as they are positive values found for a blank material. All the other laboratories in the study identified the blank materials below the working range of the test.

The valid data from the contaminated test materials (D, E, K, L, and N) were then analyzed for identification of outliers applying Cochran and Grubbs tests (20). Results from Laboratory L could not be included in the statistical analysis, as only one value per material was available.

The Cochran test showed Laboratory G for Material K and Laboratory P for Material E as outliers. This test was applied again after these outliers were removed. Laboratory J for Material E was also excluded in a second round. The Grubbs single and pair values tests were then applied; no further outliers were identified.

The mean values assigned for OA-toxins for the test materials were 98.8, 175.4, 242.8, 255.0, and 275.0 µg total equivalents OA/kg for Materials E, L, D, N, and K, respectively (Table 7).

Values obtained for repeatability SD (S_r) ranged from 7.3 µg/kg for Material E to 19.6 µg/kg for Material L, with repeatability RSDs (RSD_r) from 5.4% for Material K to 11.2% for Material L (Table 7). The reproducibility SD (S_R) calculated for the five test materials ranged from 10.7 to 23.2 µg/kg, with reproducibility RSD (RSD_R) values from 7.6 to 13.2% for Materials K and L, respectively (Table 7).

The HorRat values obtained were 0.4 for Materials D, K, and N, 0.5 for Material E, and 0.6 for Material L (Table 7), indicating a very good performance of the method. These values are just at the lower limit of the range considered as normally expected for a good reproducibility of a method (0.5 < HorRat ≤ 1.5), according to the AOAC guidelines (20). HorRat values between 0.64 and 2.61 for OA-toxins group (21), 0.3 and 2.0 for paralytic

Table 8. Results from the recovery experiment carried out during Day 2 of the interlaboratory study

Lab code	µg OA total eq./kg			Recovery, %
	BM ^a	Spiked concn.	BM + OA ^b	
A	—	161	172	107.1
B	—	161	162	100.7
C	—	161	155	96.3
D	—	161	115	71.6
E	—	161	124	77.3
F	—	161	138	85.5
G	—	161	162	100.7
H	—	161	131	81.1
I	—	161	152	94.4
J	—	161	197	122.3
K	—	161	152	94.4
L	—	161	196	121.6
M	—	161	153	95.0
N	—	161	174	108.3
O	—	161	155	96.3
P	—	161	185	114.7
Mean recovery, %				98.0
SD				14.2
RSD, %				14.5

^a BM = Blank material. No OA toxins were detected; therefore, a concentration of zero was considered for calculation purposes.

^b BM + OA = Concentration of the samples spiked with 161 µg/kg.

shellfish toxins (22) and 1.1 to 2.4 for domoic acid (23) were previously described for other methods.

The statistical analysis was also carried out including outliers (Table 7). Although there were some differences when including outlier values, repeatability and reproducibility remained satisfactory and within the expected values for this type of interlaboratory study.

Although the main objective of the validation study was to determine the repeatability and between-laboratory reproducibility of the OkaTest kit, a recovery assay was also carried out. A scallop blank sample (BM) was spiked with OA by each laboratory, and the recovery of OkaTest calculated. Recovery values from all participants ranged from 71.6 to 122.3%. The mean and RSD were 98.0 and 14.5%, respectively (Table 8). These recoveries met the criteria set in the AOAC Guidelines for Single Laboratory Validation of Chemical Methods for Dietary Supplements and Botanicals (24).

Comments from Participants

Most participants reported that the SOP for the method provided all the information they needed to perform the assay and that they did not have difficulties understanding any part of it. Some comments were made about the phosphatase preparation. Those led to the conclusion that the use of a nonorbital shaker does not always guarantee full dissolution of this reagent. Manual mixing, longer preparation, and a final visual check of the solution should be included in the SOP. Other

minor comments were made, and were answered or resolved by the study director.

Conclusions

The precision and recovery values determined in this study for OkaTest can be considered satisfactory for this type of methodology and the concentration range required. The colorimetric PPIA, OkaTest, could be used as an assay complementary to the reference method for determination of the OA toxins group in molluscs according to the Commission Regulations (EC) No. 2074/2005 and No. 15/2011. Additional methods have to be implemented in a laboratory to analyze all regulated lipophilic marine biotoxins.

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ISSC Method Application and Single Lab Validation Checklist For Acceptance of a Method for Use in the NSSP

The purpose of single laboratory validation in the National Shellfish Sanitation Program (NSSP) is to ensure that the analytical method under consideration for adoption by the NSSP is fit for its intended use in the Program. A Checklist has been developed which explores and articulates the need for the method in the NSSP; provides an itemized list of method documentation requirements; and, sets forth the performance characteristics to be tested as part of the overall process of single laboratory validation. For ease in application, the performance characteristics listed under validation criteria on the Checklist have been defined and accompany the Checklist as part of the process of single laboratory validation. Further a generic protocol has been developed that provides the basic framework for integrating the requirements for the single laboratory validation of all analytical methods intended for adoption by the NSSP. Methods submitted to the Interstate Shellfish Sanitation Conference (ISSC) Laboratory Methods Review (LMR) Committee for acceptance will require, at a minimum, six (6) months for review from the date of submission.

Name of the New Method		Receptor Binding Assay (RBA) for Paralytic Shellfish Poisoning (PSP) Toxicity Determination	
Name of the Method Developer		Dr. Fran Van Dolah	
Developer Contact Information		Tel: (843) 725-4864 Email: Fran.vandolah@noaa.gov	
Checklist	Y/N	Submitter Comments	
A. Need for the New Method			
1. Clearly define the need for which the method has been developed.	Y	<p>Paralytic shellfish poisoning (PSP) is the human intoxication that results from the consumption of seafood, primarily bivalve molluscs, contaminated with natural, algal-derived toxins known as paralytic shellfish toxins (PSTs) or the saxitoxins (STXs). This family of neurotoxins binds to voltage-gated sodium channels, thereby attenuating action potentials by preventing the passage of sodium ions across the membrane. Symptoms include tingling, numbness, headaches, weakness, and difficulty breathing. Medical treatment is to provide respiratory support, without which the prognosis can be fatal. To protect human health, seafood harvesting bans are implemented when toxins exceed a safe guidance level (80 µg STX equivalents per 100 g tissue or 800 µg STX equivalents per kg). Successful monitoring and management programs are attributed with minimizing the number of PSP cases and associated deaths.</p> <p>The mouse bioassay (MBA) has long-served as the gold standard method for detecting PSP in regulatory environments. Even though the MBA is an NSSP Approved Method for Marine Biotxin Testing, there are numerous reasons for considering alternative methods for PSP detection. Disadvantages of the MBA include high variability and the use of live animals. Given these limitations of the MBA, particularly the ethical concerns of using live animals, there have been great strides in method development and validation for alternative approaches.</p> <p>Recently, the post-column oxidation liquid chromatographic method (PCOX) for PSP detection was accepted as an NSSP Approved Limited Use Method, providing an alternative to the MBA. While some laboratories are in the process of transitioning to this</p>	

		<p>method, implementation requires costly instrumentation and skilled personnel. Furthermore, the PCOX method identifies and quantifies individual PSP toxins. Toxicity equivalency factors must then be taken into consideration to calculate the expected overall toxicity in µg STX equivalents per 100 g tissue.</p> <p>The proposed receptor binding assay (RBA) addresses the major shortcomings of the PCOX and MBA by quantitatively measuring the overall PSP toxicity and doing so without the need of live animals, respectively. The RBA relies on the interaction of the toxins with the native receptor site (i.e., voltage-gated sodium channels). In this functional assay toxins bind to their receptors according to their affinity, yielding an integrated toxic potency. The RBA is more sensitive than the MBA, allowing monitoring laboratories earlier warning capabilities as toxins become elevated. The RBA has successfully undergone AOAC single laboratory validation (Van Dolah et al. 2009 - Appendix II) and a full collaborative study (Van Dolah et al. 2012 - Appendix III). The RBA is now considered an AOAC Official Method of Analysis (OMA 2011.27 - Appendix IV). This proposal provides data from the AOAC studies as well as additional data to seek consideration for the RBA to be an NSSP Approved Limited Use Method.</p>
2. What is the intended purpose of the method?	Y	<p>This method is intended for use as an NSSP Approved Limited Use Method for screening for PSP toxicity in shellfish. Applications include: (1) Growing Area Survey & Classification and (2) Controlled Relaying. The RBA serves as an alternative to the MBA in these applications, offering a measure of integrated toxicity with high throughput and the elimination of live animal testing.</p>
3. Is there an acknowledged need for this method in the NSSP?	Y	<p>Yes, there is an acknowledged need for this method in the NSSP. Even though the MBA and PCOX methods have been respectively NSSP Approved and Approved for Limited Use, there remains a need for the proposed method. The RBA would provide an alternative to (1) the MBA, which uses live animals, and (2) the PCOX method, which requires costly equipment and skilled personnel and offers low throughput.</p>
4. What type of method? i.e. chemical, molecular, culture, etc.	Y	<p>Molecular. The RBA is a functional assay, whereby toxins present in the standard/sample bind to sodium channel preparations in the assay. Radiolabeled toxins are added to solution to compete with toxins present in the standard/sample for binding sites, and thus a decrease in signal from radiolabeled toxins represents an increase in standard/sample toxicity. This competitive RBA allows for quantitation that directly relates to the composite toxicity of the sample.</p>
B. Method Documentation		
1. Method documentation includes the following information:		
Method Title	Y	Receptor Binding Assay (RBA) for Paralytic Shellfish Poisoning (PSP) Toxicity Determination
Method Scope	Y	The RBA provides a high throughput, sensitive, accurate, quantitative assay for PSP toxins in shellfish. The method is being submitted for consideration as an NSSP Approved Limited Use Method for the purposes of screening for PSP toxicity.

References	Y	<p>Van Dolah et al. 2009. Single-laboratory validation of the microplate receptor binding assay for paralytic shellfish toxins in shellfish. <i>Journal of AOAC International</i> 92(6): 1705-1713. See Appendix II.</p> <p>Van Dolah et al. 2012. Determination of paralytic shellfish poisoning toxins in shellfish by receptor binding assay: Collaborative study. <i>Journal of AOAC International</i> 95(3): 795-812. See Appendix III.</p> <p>OMA 2011.27. AOAC Official Method 2011.27 Paralytic shellfish toxins (PSTs) in shellfish, receptor binding assay. In <i>Official Methods of Analysis of AOAC International</i>. http://www.eoma.aoac.org. See Appendix IV.</p>
Principle	Y	<p>This assay is based on the interaction between the toxins and their native receptor, the voltage-gated sodium channels. All PSTs bind to site 1 of the voltage-gated sodium channels according to their potency, resulting in a measure of integrated potency (independent of knowing which toxin congeners are present) similar to mouse intraperitoneal potency. In the RBA, tritiated saxitoxin (³H-STX) competes with unlabeled PSTs in the homogenized and extracted shellfish sample for a finite number of available receptor sites in a rat brain membrane preparation. After a binding equilibrium is reached, unbound ³H-STX is removed by filtration and the remaining ³H-STX is measured with a scintillation counter (as counts per minute or CPM). The amount of ³H-STX present is indirectly related to the amount of unlabeled PSTs in the sample. Scintillation counting can be conducted using traditional scintillation counters or microplate counting. However, the microplate format is preferred as it minimizes sample handling and the amount of radioactivity used.</p>
Any Proprietary Aspects	N	None. All reagents can be prepared or purchased.
Equipment Required	Y	<p>The following list identifies the equipment and supplies needed for conducting the RBA.</p> <p>For the assay:</p> <ul style="list-style-type: none"> (a) Scintillation counter (traditional or microplate) (b) An 8-channel pipettor (5-200 µl variable volume and disposable tips) (c) Micropipettors (1-1000 µl variable volumes and disposable tips) (d) 96-well microtitre filter plate (1 µm pore size type GF/B glass fiber filter/0.65 µm pore size Durapore support membrane (Millipore, Bedford, MA; Cat. No. MSFB N6B 50) (e) MultiScreen vacuum manifold (Millipore; Cat. No. NSVMHTS00) (f) Vacuum pump (g) Centrifuge tubes (15 and 50 ml, conical, plastic) (h) Mini dilution tubes in 96-tube array (i) Reagent reservoirs (j) Ice bucket and ice (k) Vortex mixer (l) Sealing tape (Millipore; Cat. No. MATA HCL00) (m) Volumetric flask or graduated beaker (1 L) (n) -80 °C freezer (o) Refrigerator

		<p>Additional supplies when using a traditional scintillation counter (as opposed to a microplate counter):</p> <p>(p) MultiScreen punch device (Millipore; Cat. No. MAMP 096 08)</p> <p>(q) MultiScreen disposable punch tips (Millipore; Cat. No. MADP 196 10)</p> <p>(r) MultiScreen punch kit B for 4 ml vials (Millipore; Cat. No. MAPK 896 0B)</p> <p>(s) Scintillation vials (4 ml)</p> <p>For sample extraction:</p> <p>(t) Blender or homogenizer for sample homogenization</p> <p>(u) Pipets</p> <p>(v) Centrifuge tubes (15 ml, conical, plastic)</p> <p>(w) pH meter or pH paper</p> <p>(x) Hot plate or water bath</p> <p>(y) Graduated centrifuge tubes (15 ml)</p> <p>(z) Centrifuge and rotor for 15 ml tubes</p> <p>For rat brain isolation:</p> <p>(aa) Teflon/glass homogenizer (Motorized tapered Teflon pestle and glass tube (15 ml)</p> <p>(bb) Motorized tissue homogenizer (Polytron or small handheld blender)</p> <p>(cc) High-speed centrifuge and fixed angle rotor (20 000 x g rcf)</p> <p>(dd) Centrifuge tubes (12-15 ml, rated for 20 000 x g)</p> <p>(ee) plastic cryovials (2 ml)</p> <p>(ff) Graduated beaker (300 or 500 ml)</p> <p>(hh) Pipets (5-10 ml, disposable)</p> <p>(ii) Forceps</p> <p>(jj) Ice bucket and ice</p> <p>(kk) top loading balance</p>
<p>Reagents Required</p>	<p>Y</p>	<p>For the assay:</p> <p>(a) STX diHCl standards (NIST RM 8642; available through the National Institute of Standards and Technology; www.nist.gov) [This is the same standard used for the MBA]</p> <p>(b) ³H-STX (0.1 mCi per ml, ≥10 Ci per mmol, ≥90% radiochemical purity; available through American Radiolabeled Chemicals, St. Louis, MO)</p> <p>(c) 3-Morpholinopropanesulfonic acid (MOPS; Sigma; St. Louis, MO; Cat. No. M3183-500G [or equivalent])</p> <p>(d) Choline chloride (Sigma; Cat. No. C7527-500G [or equivalent])</p> <p>For microplate counter only:</p> <p>(e) Ultima Gold liquid scintillation cocktail (PerkinElmer Inc.; Waltham, MA; Cat. No. 6013321 [or equivalent])</p> <p>For traditional counter only:</p> <p>(f) Scintiverse BD liquid scintillation cocktail (Fisher Scientific; Waltham, MA; Cat. No. SX-18 [or equivalent])</p> <p>For sample extraction:</p> <p>(g) Hydrochloric acid (HCl; 1.0 and 0.1 M)</p> <p>(h) Sodium hydroxide (0.1 M)</p> <p>(i) Water (distilled or deionized [18 μΩ])</p> <p>For rat brain isolation:</p> <p>(j) 20 rat brains (male, 6-week old Sprague-Dawley;</p>

		<p>available through Hilltop Lab Animals, Inc., Scottsdale, PA; www.hilltoplabs.com [or equivalent]) (k) MOPS, pH 7.4 (Sigma, St. Louis, MO; Cat. No. M3183-500G [or equivalent]) (l) Choline chloride (100 mM; Sigma; Cat. No. C7527-500G [or equivalent]) (m) Phenyl methylsulfonyl fluoride (PMSF; Sigma, St. Louis, MO; Cat. No. P7626) (n) Isopropanol (o) Micro bicinchoninic acid (BCA) protein assay (Pierce, Rockford, IL)</p>
Sample Collection, Preservation and Storage Requirements	Y	<p>A representative shellfish sample should include 12 market size organisms pooled together (should be at least 100 g). Clean the outside of shellfish with running tap water. Open the shell by cutting into the adductor muscle, being careful to not cut or damage the viscera. Rinse the inside to remove sand and dirt and remove tissue from ~12 organisms. Collect the tissue on a number 10 sieve and allow to drain for ~5 minutes. Remove any obvious pieces of shell or debris. Transfer meat to blender or homogenizer and blend until homogeneous. This homogenate is then extracted for toxins. For the detailed sample extraction procedure see Sample Extraction in Appendix A. Shellfish homogenates must be tested immediately or stored frozen prior to analysis. Saxitoxin standards must be stored refrigerated and ³H-STX must be stored at -80 °C. The rat brain preparation can be produced in bulk, partitioned into aliquots, and stored long-term at -80 °C until use.</p>
Safety Requirements	Y	<p>General safety requirements (e.g., personal protective equipment including gloves, safety glasses, and laboratory coat) for working with toxins, biological reagents, and radioactive material must be followed. Users must be trained in and follow all in-house safety procedures for working with toxins and radiolabeled materials. Even though low levels of radiation are used for this assay, users must follow all local, state and federal laws and procedures regarding the receipt, use, and disposal of isotopes. Please see Appendix C for further safety requirements.</p>
Clear and Easy to Follow Step-by-Step Procedure	Y	<p>The protocol is very clear and easy to follow. Please see the detailed protocol below in Appendix A.</p>
Quality Control Steps Specific for this Method	Y	<p>Quality control steps are in place to determine if assay results are acceptable:</p> <p>(a) The slope of the standard curve must be between -0.8 and -1.2 (theoretical slope is -1). If the slope of a standard curve from a given assay falls outside of this range, the data should be considered unacceptable and the assay must be rerun.</p> <p>(b) The RSDs of triplicate counts per minute (CPMs) for the standards must be below 30%.</p> <p>(c) If the IC₅₀ (inhibitory concentration at which CPM is 50% max) is out of the acceptable range (2.0 nM ± 30%), the data should be considered unacceptable and the assay should be rerun.</p> <p>(d) A QC sample should always be included and found to be in range. Typically a 1.8 x 10⁻⁸ M STX concentration</p>

		<p>(3 nM STX in-well concentration) is run as a QC and should be within 30%. Results outside of this range should trigger consideration of assay acceptance.</p> <p>The following criteria must be met to accept sample measurement:</p> <p>(e) For sample measurement, quantitation should only be done on sample dilutions that fall within the linear range. As such, binding (B, measured as counts per minute) scaled by the maximum binding (B₀) should be between 0.2-0.7 for sample quantitation to be performed (any sample falling outside of this range is considered out of the dynamic range). If B/B₀ > 0.7, the concentration is too low to be quantified and should be reported as below the limit of detection (LOD). If B/B₀ < 0.2, the sample should be diluted and rerun if quantitation is needed.</p> <p>(f) The RSDs for the sample CPMs should be ≤ 30%.</p> <p>These quality control criteria are also stated in section H in Appendix IV.</p>
C. Validation Criteria		
<p>1. Accuracy / Trueness</p>	<p>Y</p>	<p>Validation data presented in Section C are from both the SLV (Van Dolah et al. 2009) and the collaborative study (Van Dolah et al. 2012). Nine laboratories from six countries completed the collaborative study. There were a total of 21 shellfish homogenates tested in three different assays on independent days. Different shellfish species from a range of geographical locations were used in the study: blue mussel (<i>Mytilus edulis</i>) from the U.S. east and west coasts, California mussel (<i>Mytilus californianus</i>) from the U.S. west coast, chorito mussel (<i>Mytilus chilensis</i>) from Chile, green mussel (<i>Perna canaliculus</i>) from New Zealand, Atlantic surfclam (<i>Spisula solidissima</i>) from the U.S. east coast, butter clam (<i>Saxidomus gigantea</i>) from the U.S. west coast, almeja clam (<i>Venus antiqua</i>) from Chile, and Atlantic sea scallop (<i>Placopecten magellanicus</i>) from the U.S. east coast. Samples included those that were naturally contaminated, those that were spiked, and another that served as a negative control.</p> <p>Accuracy was evaluated based on recovery. As also stated under Section C. 4., Recovery of the QC check sample (3 nM in-well solution) was 99.3% (Appendix II).</p> <p>During the SLV recovery was evaluated for STX standard spiked into mussel tissue at concentrations below, at and above the regulatory guidance level. Recovery for the nominal spike at 40 µg STX eq 100 g⁻¹ was 115%. At 80 µg STX eq 100 g⁻¹, recovery was found to be 129%. At a nominal spike of 120 µg STX eq 100 g⁻¹, recovery was 121% (Appendix II).</p> <p>During the collaborative study, recovery of PSTs from shellfish was found to be 84.4% (when spiked with 20 µg STX eq 100 g⁻¹), 93.3% (when spiked with 50 µg STX eq 100 g⁻¹), and 88.1% (when spiked with 120 µg STX eq 100 g⁻¹). See Appendix III.</p>
<p>2. Measurement Uncertainty</p>	<p>Y</p>	<p>ND</p>

<p>3. Precision Characteristics (repeatability and reproducibility)</p>	<p>Y</p>	<p>Repeatability (RSD_r) was determined during the SLV on six naturally contaminated shellfish samples on five independent days and was found to be 17.7%. See Appendix II.</p> <p>The reproducibility (RSD_R) during the collaborative study was found to be 33.2% for all laboratories. However, upon removing the results from the one laboratory that had no previous RBA experience, the RSD_R was 28.7%. If data from routine users of the RBA were evaluated, the RSD_R was 23.1%. See Appendix III.</p> <p>Repeatability (RSD_r) during the collaborative study ranged from 11.8-34.4%. For routine users of the RBA, the average $RSD_r = 17.1%$, consistent with the RSD_r obtained during the SLV. See Appendix III.</p>
<p>4. Recovery</p>	<p>Y</p>	<p>Recovery of the QC check sample (3 nM in-well solution) was 99.3% (Appendix II).</p> <p>During the SLV recovery was evaluated for STX standard spiked into mussel tissue at concentrations below, at and above the regulatory guidance level. Recovery for the nominal spike at $40 \mu\text{g STX eq } 100 \text{ g}^{-1}$ was 115%. At $80 \mu\text{g STX eq } 100 \text{ g}^{-1}$, recovery was found to be 129%. At a nominal spike of $120 \mu\text{g STX eq } 100 \text{ g}^{-1}$, recovery was 121% (Appendix II).</p> <p>During the collaborative study, recovery of PSTs from shellfish was found to be 84.4% (when spiked with $20 \mu\text{g STX eq } 100 \text{ g}^{-1}$), 93.3% (when spiked with $50 \mu\text{g STX eq } 100 \text{ g}^{-1}$), and 88.1% (when spiked with $120 \mu\text{g STX eq } 100 \text{ g}^{-1}$). See Appendix III.</p>
<p>5. Specificity</p>	<p>Y</p>	<p>The RBA is specific to toxins that bind to site 1 of voltage-gated sodium channels. This includes all PSP congeners, whereby binding affinity is proportional to potency. Tetrodotoxin also binds to site 1 of the sodium channels, yet the typical combinations of sources, vectors, and geographical regions of tetrodotoxin and the saxitoxins differ.</p>
<p>6. Working and Linear Ranges</p>	<p>Y</p>	<p>The dynamic range of the assay was determined to be 1.2-10.0 nM in-well concentration (Appendix II). Linearity assessment was conducted with three calibration standards (1.5, 3.0, and 6.0 nM STX in -well concentration) on five independent days. The linear regression yielded a slope of 0.98 and an $r^2 = 0.97$ (Appendix II).</p> <p>During the collaborative study, the assay was set for the critical range of shellfish toxicities below, near and just above the regulatory guidance level ($\sim 15\text{-}240 \mu\text{g STX eq } 100 \text{ g}^{-1}$ or $\sim 150\text{-}2400 \mu\text{g STX eq kg}^{-1}$). Appendix III.</p>
<p>7. Limit of Detection</p>	<p>Y</p>	<p>The LOD, as determined in the collaborative study, is $4.5 \mu\text{g STX eq } 100 \text{ g}^{-1}$ or $45 \mu\text{g STX eq kg}^{-1}$ See Appendix III.</p>
<p>8. Limit of Quantitation / Sensitivity</p>	<p>Y</p>	<p>The limit of quantitation (LOQ) was empirically determined as the concentration in a 10-fold diluted sample that resulted in a B/B0 of 0.7 (more conservative than the 0.8 typically used as the cut off for such assays). The LOQ was determined to be $5.3 \mu\text{g STX eq } 100 \text{ g}^{-1}$ during the SLV (Appendix II).</p>

		The LOQ of the RBA is 12.6 µg STX eq 100 g ⁻¹ or 126 µg STX eq kg ⁻¹ , as compared to the MBA LOQ of ~40 µg STX eq 100 g ⁻¹ (or ~400 µg STX eq kg ⁻¹). See Appendix III.
9. Ruggedness	Y	<p>Ruggedness was addressed and critical steps were noted that could affect precision and accuracy. It was deemed important to clarify the shellfish extracts by centrifugation prior to performing the assay, particularly if the sample was refrigerated or frozen. The rat brain preparations should be vortexed frequently to ensure the synaptosomes are in suspension, and the buffer should be ice cold to ensure that toxins are not released from the receptor. Assay plate filtration should be at a rate of 2-5 seconds. Lastly, a minimum of 30 minutes should be allowed before reading the plates after scintillation liquid is added such that scintillant can penetrate the filters.</p> <p>For more detail please refer to Appendix II and Appendix III.</p>
10. Matrix Effects	Y	No matrix effects were reported. Minimum dilutions of shellfish extracts were 10-fold and were found to be sufficient to eliminate matrix effects. See Appendix III.
11. Comparability (if intended as a substitute for an established method accepted by the NSSP)	Y	<p>The RBA was compared to the MBA and the pre-column oxidation (Pre-COX) liquid chromatography with fluorescence detection (LC-FD) approach during the SLV.</p> <p>RBA results compared well to those obtained by the MBA in two separate studies. In one component of the SLV, six naturally contaminated samples (clams, mussels, and sea scallops) were tested by RBA and MBA. Between-assay RSDs ranged from 9 to 25% (mean 17.7%). An $r^2 = 0.98$ was obtained, with a slope of 1.29. In the second component of the SLV, which included 110 naturally contaminated shellfish, an $r^2 = 0.88$ and a slope of 1.32 was obtained (Appendix II).</p> <p>Nine naturally contaminated samples (six blue mussels and three scallops) were extracted and analyzed by RBA and Pre-COX. Samples were analyzed using the RBA following the typical extraction (0.1 M HCl), but also following the extraction procedure used for the Pre-COX method (1% acetic acid). A good correlation was found between the two methods for both extraction methods. When the RBA samples were extracted with HCl, the RBA compared to the Pre-COX yielded an $r^2 = 0.98$ and a slope of 1.39. When samples were extracted the same for both methods (acetic acid), the correlation was slightly improved with an $r^2 = 0.99$ and a slope of 1.32 (Appendix II).</p> <p>During the collaborative study, ten laboratories from seven countries performed the RBA. Additionally three of the laboratories conducted the MBA, and one laboratory tested the samples using the Pre-COX LC-FD. The MBA and RBA data comparison yielded an $r^2 = 0.84$ and a slope of 1.63. The LC-FD and RBA data comparison yielded an $r^2 = 0.92$ and a slope of 1.20. Both RBA and LC-FD methods generally report higher toxicity in shellfish, especially at or near the guidance level, relative to the MBA. This provides a conservative measure and allows for an earlier warning of developing</p>

		toxicity. See Appendix III.
D. Other Information		
1. Cost of the Method	Y	The estimated cost per 96-well plate assay is ~\$95.00. Including standards and samples with triplicate measurements (as well as three dilutions per sample [ranging from 3.5-600 µg STX eq 100 g ⁻¹] to ensure the unknown samples fall within linear range of assay), the cost per sample for quantitation would be ~\$13.60. If running multiple plates or in screening mode, sample costs would be reduced.
2. Special Technical Skills Required to Perform the Method	Y	General laboratory training is necessary (this would include being able to prepare reagent solutions, pipetting, centrifugation, and simple calculations). Additional training for working with low levels of radioactive material is required.
3. Special Equipment Required and Associated Cost	Y	A microplate scintillation counter is needed and the cost is ~\$60-100K for a new counter, depending on the brand and number of simultaneous detectors. However, used instruments can be purchased for ~\$13K.
4. Abbreviations and Acronyms Defined	Y	A list of abbreviations and acronyms is provided below in Appendix I.
5. Details of Turn Around Times (time involved to complete the method)	Y	Microplate scintillation counting provides the ability to test multiple samples simultaneously with a turn around time for data in approximately 3 hours. Up to six plates per analyst are possible in one day, yielding a throughput of 42 samples per day.
6. Provide Brief Overview of the Quality Systems Used in the Lab	Y	The Center for Food Safety and Applied Nutrition (CFSAN) Quality System (QS) provides guidance to (1) design and develop processes, products, and services related to CFSAN's mission, the FDA's regulatory mission, and critical management and administrative support services, and (2) continually improve and strengthen product and service quality. The Laboratory Quality Assurance program serves as CFSAN's logical application of QS to Center laboratories and lab-based activities. The third edition (October 2009) of the Laboratory Quality Manual was followed. Standard reference materials for saxitoxin are obtained through the National Institute of Standards and Technology (NIST) and are accompanied by a Report of Investigation (See Appendix V). The standard reference saxitoxin used in the RBA is the same as that employed with the MBA. The 3H-STX is obtained through American Radiolabeled Chemicals, Inc., and is accompanied by a Technical Data Sheet with lot specifications (Appendix VI).
Submitters Signature	Date:	
Submission of Validation Data and Draft Method to Committee	Date:	
Reviewing Members	Date:	

Accepted	Date:
Recommendations for Further Work	Date:
Comments:	

DEFINITIONS

1. **Accuracy/Trueness** - Closeness of agreement between a test result and the accepted reference value.
2. **Analyte/measurand** - The specific organism or chemical substance sought or determined in a sample.
3. **Blank** - Sample material containing no detectable level of the analyte or measurand of interest that is subjected to the analytical process and monitors contamination during analysis.
4. **Comparability** - The acceptability of a new or modified method as a substitute for an established method in the NSSP. Comparability must be demonstrated for each substrate or tissue type by season and geographic area if applicable.
5. **Fit for purpose** - The analytical method is appropriate to the purpose for which the results are likely to be used.
6. **HORRAT value** - HORRAT values give a measure of the acceptability of the precision characteristics of a method.⁴
7. **Limit of Detection** - the minimum concentration at which the analyte or measurand can be identified. Limit of detection is matrix and analyte/measurand dependent.⁴
8. **Limit of Quantitation/Sensitivity** - the minimum concentration of the analyte or measurand that can be quantified with an acceptable level of precision and accuracy under the conditions of the test.
9. **Linear Range** - the range within the working range where the results are proportional to the concentration of the analyte or measurand present in the sample.
10. **Measurement Uncertainty** - A single parameter (usually a standard deviation or confidence interval) expressing the possible range of values around the measured result within which the true value is expected to be with a stated degree of probability. It takes into account all recognized effects operating on the result including: overall precision of the complete method, the method and laboratory bias and matrix effects.
11. **Matrix** - The component or substrate of a test sample.
12. **Method Validation** - The process of verifying that a method is fit for purpose.¹
13. **Precision** - the closeness of agreement between independent test results obtained under stipulated conditions.^{1, 2}
 There are two components of precision:
 - a. **Repeatability** - the measure of agreement of replicate tests carried out on the same sample in the same laboratory by the same analyst within short intervals of time.
 - b. **Reproducibility** - the measure of agreement between tests carried out in different laboratories. In single laboratory validation studies reproducibility is the closeness of agreement between results obtained with the same method on replicate analytical portions with different analysts or with the same analyst on different days.

14. **Quality System** - The laboratory's quality system is the process by which the laboratory conducts its activities so as to provide data of known and documented quality with which to demonstrate regulatory compliance and for other decision-making purposes. This system includes a process by which appropriate analytical methods are selected, their capability is evaluated, and their performance is documented. The quality system shall be documented in the laboratory's quality manual.
15. **Recovery** - The fraction or percentage of an analyte or measurand recovered following sample analysis.
16. **Ruggedness** - the ability of a particular method to withstand relatively minor changes in analytical technique, reagents, or environmental factors likely to arise in different test environments.⁴
17. **Specificity** - the ability of a method to measure only what it is intended to measure.¹
18. **Working Range** - the range of analyte or measurand concentration over which the method is applied.

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2. IUPAC Technical Report, 2002. Harmonized Guidelines for Single-Laboratory Validation of Methods of Analysis, Pure Appl. Chem., Vol. 74, (5): 835-855.
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5. National Environmental Laboratory Accreditation. , 2003. Standards. June 5.
6. EPA. 2004. EPA Microbiological Alternate Procedure Test Procedure (ATP) Protocol for Drinking Water, Ambient Water, and Wastewater Monitoring Methods: Guidance. U.S. Environmental Protection Agency (EPA), Office of Water Engineering and Analysis Division, 1200 Pennsylvania Avenue, NW, (4303T), Washington, DC 20460. April.

Appendix A: RBA Step-by-Step Procedure

A. Sample Extraction

- a. The extraction detailed below represents a small scale MBA extraction procedure. The actual MBA extraction could be used instead of the small scale version described here.
- b. Accurately weigh 5.0 g of tissue homogenate into a tared, labeled 15 ml conical tube.
- c. Add 5.0 ml of 0.1 M HCl, vortex, and check pH.
 - i. If necessary, adjust pH to 3.0-4.0 as determined by a pH meter or pH paper. To lower pH, add 1 M HCl dropwise with mixing; to raise pH, add 0.1 M NaOH dropwise with mixing.
- d. Place the tube in a beaker of boiling water on hot plate (or in a water bath) for 5 min with the caps loosened.
- e. Remove and cool to room temperature.
- f. Check pH and, if necessary, adjust cooled mixture to 3.0-4.0 as described above.
- g. Transfer entire contents to a labeled, graduated centrifuge tube and dilute volumetrically to 10 ml.
- h. Gently stir contents to homogeneity and then allow to settle until a portion of supernatant is translucent and can be decanted free of solids.
- i. Pour 5-7 ml of the translucent supernatant into a labeled centrifuge tube.
- j. Centrifuge at 3000 x *g* for 10 min.
- k. Retain clarified supernatant and transfer to a clean, labeled centrifuge tube.
- l. Store extracts at -20 °C until tested in RBA.

B. Preparation of Stock Solutions and Standards

- a. Assay buffer: 100 mM MOPS/100 mM choline chloride, pH 7.4
 - i. Weigh 20.9 g MOPS and 13.96 g choline chloride and add to 900 ml distilled or milli-Q water.
 - ii. Adjust pH to 7.4 with NaOH while stirring.
 - iii. Bring to a final volume of 1 L with distilled or milli-Q water.
 - iv. Store at 4 °C.
- b. Radioligand solution: ³H-STX
 - i. Calculate the concentration of ³H-STX stock provided by the supplier. Suppliers generally provide specific activity in Ci/mmol (~10-30 Ci/mmol) and activity in mCi/ml (~0.05-0.1 mCi/ml), from which the molar concentration can be calculated.
 - ii. Prepare 4 ml of a 15 nM working stock of ³H-STX fresh daily in 100 mM MOPS/100 mM choline chloride buffer. This will provide sufficient volume for one 96-well plate.
 - iii. Measure total counts of each working stock prior to running an assay. Add 36 µl of working stock ³H-STX in buffer to a liquid scintillation counter vial with 4 ml scintillant and count on a traditional liquid scintillation counter to confirm correct dilution. The CPM should be consistent and within 15% of expected value.

- c. Unlabeled STX standard working solution: The STX diHCl standard (NIST RM 8642 STX diHCl) is provided at a concentration of 268.8 μM (100 $\mu\text{g}/\text{ml}$).
- i. A bulk standard curve can be made up in advance and stored at 4 °C for up to one month. The use of a bulk standard curve minimizes time needed for routine analyses and improves repeatability.
 - ii. Make up 3 mM HCl (e.g., from a 3 M stock, 50 μl in 50 ml) and use for the serial dilutions.
 - iii. Serial dilutions should result in the following stock concentrations (M):
 1. 6×10^{-6} [100 μl 268.8 μM STX + 4.38 ml 0.003 M HCl]
 2. 6×10^{-7} [500 μl 6×10^{-6} M STX + 4.5 ml 0.003 M HCl]
 3. 1.8×10^{-7} [1.5 ml 6×10^{-7} M STX + 3.5 ml 0.003 M HCl]
 4. 6×10^{-8} [500 μl 6×10^{-7} M STX + 4.5 ml 0.003 M HCl]
 5. 1.8×10^{-8} [500 μl 1.8×10^{-7} M STX + 4.5 ml 0.003 M HCl]
 6. 6×10^{-9} [500 μl 6×10^{-8} M STX + 4.5 ml 0.003 M HCl]
 7. 6×10^{-10} [500 μl 6×10^{-9} M STX + 4.5 ml 0.003 M HCl]
 8. 5 ml 0.003 M HCl.
- d. Interassay calibration standard (QC check): Reference standard STX (1.8×10^{-8} M STX) in 3 mM HCl. For long-term storage keep at -80 °C; for routine use (up to one month), store at 4 °C.
- e. Rat brain membrane preparation: Prepare bulk rat brain membrane preparations (Appendix B) and store at -80 °C.
- i. Thaw an aliquot of rat brain preparation on ice.
 - ii. Dilute membrane preparation with cold (4 °C) 100 mM MOPS/100 mM choline chloride, pH 7.4 to yield a working stock with a protein concentration of 1.0 mg/ml.
 - iii. Vortex vigorously to achieve a visibly homogeneous suspension.
 - iv. Keep the diluted membrane preparation on ice.
- C. Performing the Assay
- a. Plate setup: When possible use a multichannel pipet to minimize effort and increase consistency.
 - i. Run standards, samples, and QC check in triplicate.
 - ii. For quantitation, multiple dilutions per extract should be analyzed in order to obtain a value that falls within the dynamic range of the assay. A minimum sample extract dilution of 1:10 is recommended to minimize potential matrix effects.
 - iii. Use of a standard plate layout (Figure 1) is recommended. This will improve ease of analysis and can help maximize the number of samples/standards that can be analyzed per plate.
 - b. Addition of samples/standards: Add in the following order to each well-
 - i. 35 μl assay buffer
 - ii. 35 μl STX standard/QC check/sample extract
 - iii. 35 μl ^3H -STX
 - iv. 105 μl membrane preparation (ensure solution is homogeneous)
 - v. Cover the plate and incubate at 4 °C for 1 h.

- c. Assay filtration: Use the vacuum manifold attached to the vacuum pump with an in-line side arm flask to catch filtrate from the plate filtration process.
 - i. Set the vacuum pressure gauge on the pump or manifold to ~4-8" Hg (~135-270 millibar).
 - ii. Place the 96-well plate on the vacuum manifold.
 - iii. Fill any empty wells with 200 μ l MOPS/choline chloride buffer to ensure even vacuum pressure and filtration across the plate.
 - iv. Turn on vacuum. Optimum vacuum will pull the wells dry in 2-5 s.
 - v. With vacuum pump running, quickly rinse each well twice with 200 μ l ice cold MOPS/choline chloride buffer using a multichannel pipet. Maintain vacuum until liquid is removed.
- d. Preparation of the assay for counting: Remove the plastic bottom from the plate and blot the plate bottom once on absorbent towel.
 - i. For counting in microplate scintillation counter:
 1. Seal the bottom of a counting cassette with sealing tape.
 2. Place the microplate in the counting cassette.
 3. Add 50 μ l scintillation cocktail per well using multichannel pipet.
 4. Seal the top of the plate with sealing tape.
 5. Incubate for 30 min at room temperature.
 6. Place the plate in the scintillation counter and count for 1 min per well.
 - ii. For counting in traditional scintillation counter:
 1. Place the microplate in the MultiScreen punch system apparatus and place the disposable punch tips on top of the microplate.
 2. Punch the filters from the wells into scintillation vials and fill with 4 ml scintillation cocktail.
 3. Place caps on the vials and vortex.
 4. Allow vials to sit overnight in the dark.
 5. Count using a tritium window in a traditional scintillation counter.

D. Analysis of Data

- a. Curve fitting: Perform curve fitting using a four-parameter logistic fit (sigmoidal dose response curve with variable slope).
 - i. $y = \min + (\max - \min) / (1 + 10^{(x - \log IC_{50}) \text{Hill slope}})$
 - ii. where max is the top plateau representing maximum binding in CPM in the absence of competing nonradiolabeled STX (also known as B_0); min is the bottom plateau, equal to nonspecific binding in CPM in the presence of saturating nonradiolabeled STX; IC_{50} is the inhibitory concentration at which CPM are 50% of max-min); Hill slope is the slope of the curve; x axis is the log concentration of STX; and y axis is the total ligand binding in CPM (B/B_0).
- b. Sample quantification: Sample quantification is only carried out on dilutions having a B/B_0 in the range of 0.2-0.7.
 - i. Where B represents the bound $^3\text{H-STX}$ in CPM in the sample and B_0 represents the max bound $^3\text{H-STX}$ in the sample.

- ii. Sample concentration is calculated in μg STX diHCl equivalents (eq)/kg shellfish as described below:

$$\begin{aligned} & (\text{nM STX eq}) \times (\text{sample dilution}) \times [(210 \mu\text{l total volume})/35 \mu\text{l sample}] \\ & = \text{nM STX eq in extract} \end{aligned}$$

$$\begin{aligned} & (\text{nM STX diHCl eq in extract}) \times (1 \text{ L}/1000 \text{ ml}) \times (372 \text{ ng/nmol}) \times (1 \mu\text{g}/1000 \text{ ng}) \\ & = \text{g STX diHCl eq/ml} \end{aligned}$$

$$\begin{aligned} & \mu\text{g STX diHCl eq/ml} \times (\text{ml extract/g shellfish}) \times (1000\text{g}/\text{kg}) \\ & = \mu\text{g STX diHCl eq/kg} \end{aligned}$$

Figure 1. Example plate layout.

	1	2	3	4	5	6	7	8	9	10	11	12
A	10 ⁻⁶	10 ⁻⁶	10 ⁻⁶	QC	QC	QC	U3 1:50	U3 1:50	U3 1:50	U6 1:10	U6 1:10	U6 1:10
B	10 ⁻⁷	10 ⁻⁷	10 ⁻⁷	U1 1:10	U1 1:10	U1 1:10	U3 1:200	U3 1:200	U3 1:200	U6 1:50	U6 1:50	U6 1:50
C	3 x 10 ⁻⁸	3 x 10 ⁻⁸	3 x 10 ⁻⁸	U1 1:50	U1 1:50	U1 1:50	U4 1:10	U4 1:10	U4 1:10	U6 1:200	U6 1:200	U6 1:200
D	10 ⁻⁸	10 ⁻⁸	10 ⁻⁸	U1 1:200	U1 1:200	U1 1:200	U4 1:50	U4 1:50	U4 1:50	U7 1:10	U7 1:10	U7 1:10
E	3 x 10 ⁻⁹	3 x 10 ⁻⁹	3 x 10 ⁻⁹	U2 1:10	U2 1:10	U2 1:10	U4 1:200	U4 1:200	U4 1:200	U7 1:50	U7 1:50	U7 1:50
F	10 ⁻⁹	10 ⁻⁹	10 ⁻⁹	U2 1:50	U2 1:50	U2 1:50	U5 1:10	U5 1:10	U5 1:10	U7 1:200	U7 1:200	U7 1:200
G	10 ⁻¹⁰	10 ⁻¹⁰	10 ⁻¹⁰	U2 1:200	U2 1:200	U2 1:200	U5 1:50	U5 1:50	U5 1:50			
H	REF	REF	REF	U3 1:10	U3 1:10	U3 1:10	U5 1:200	U5 1:200	U5 1:200			

Concentrations indicate those of the STX standard curve; REF = reference; QC = quality control; U = unknown sample (with dilutions indicated). The same standard curve made be used for additional plates run on the same day using the same reagents (i.e., 11 samples can be run on subsequent plates).

Appendix B: Rat Brain Membrane Preparations

A. Equipment/Supplies

- a. Teflon/glass homogenizer: Tapered Teflon pestle and glass tube, 15 ml
- b. Motorized tissue homogenizer: Polytron or small hand-held blender
- c. High-speed centrifuge and fixed angle rotor: capable of 20,000 x *g*
- d. Centrifuge tubes: 12-15 ml, rated for >20,000 x *g*
- e. Plastic cryovials: 2 ml
- f. Glass beaker: 300-500 ml
- g. Pipets: disposable 5 and 10 ml
- h. Forceps.

B. Reagents

- a. 20 rat brains: male, 6-week old Sprague-Dawley (Hilltop Lab Animals, Inc., Scottdale, PA) or equivalent
- b. MOPS: pH 7.4 (Sigma, St. Louis, MO; Cat. No. M3183-500G)
- c. Choline chloride: 100 mM (Sigma; Cat. No. C7527-500G)
- d. Phenyl methylsulfonyl fluoride (PMSF): (Sigma; Cat. No. P7626)
- e. Isopropanol.

C. Procedure

- a. Prepare 1 L of 100 mM MOPS, pH 7.4, containing 100 mM choline chloride (as described in Appendix A) and 0.1 mM PMSF. PMSF must first be dissolved in isopropanol: dissolve 0.174 g PMSF in 10 ml isopropanol to make 100 mM stock. Aliquot stock and store at -20 °C. Add PMSF (1/1000, 0.1 mM final concentration) to the MOPS/choline chloride buffer fresh in the day of use.
- b. Remove the medulla and cerebellum from each brain using forceps and discard. Place cerebral cortex in a small amount of ice-cold buffer and place on ice.
- c. Place one cerebral cortex in 12.5 ml MOPS/choline Cl/PMSF, pH 7.4, in glass/Teflon homogenizer. Homogenize at 70% full speed (385 rpm) with at least 10 up and down strokes and ensure there are no visible chinks remaining in the homogenate. Keep tube in ice at all times. Pour homogenized tissue into 250 ml beaker on ice and repeat procedure with remaining cortices.
- d. Transfer pooled homogenate tissue to centrifuge tubes, balance the tubes (pairwise: using ice-cold buffer to balance), and centrifuge at 20,000 x *g* for 15 min at 4 °C.
- e. Aspirate the supernatant and resuspend pellets in ice-cold MOPS/choline Cl/PMSF, using an adequate amount to fully resuspend the pellet (5-10 ml per brain).
- f. Pool resuspended membrane preparation in a small beaker. Rinse centrifuge tubes with a small amount of ice-cold buffer to recover all of the membrane preparation. Bring total volume up to 200 ml (keep on ice).
- g. Keeping the beaker on ice, polytron (or homogenize with small handheld blender) at 70% full speed for 20 s to obtain a homogeneous solution.
- h. Aliquot 2 ml per tube into cryovials. It is critical to keep the preparation well mixed while dispensing. Keep cryotubes on ice.
- i. Freeze and store at -80 °C. This preparation is stable for at least 6 months.

D. Protein Assay

- a. Determine the protein concentration of the membrane preparation using a Pierce Micro BCA Protein Assay Reagent Kit No. 23235 (microplate method) or No. 23225 (tube method) or equivalent. The above protocol should yield ~6-8 mg protein/ml of rat membrane preparation.
- b. Determine the membrane dilution needed for the assay. The protein concentration in the daily working stock should be 1 mg/ml (which yields a diluted concentration of 0.5 mg/ml in-assay concentration). Based on the protein concentration determined using the protein assay, dilute rat membrane preparation with buffer to 1 mg/ml. It is this diluted membrane preparation that is used in the assay.
- c. Protein concentrations must be determined and new dilutions calculated accordingly for each new batch of membranes prepared.

Appendix C: Radiation Safety Requirements

- A. All users must follow all local, state, and federal laws and procedures regarding receipt, use and disposal of isotopes.
- B. All users must be trained in and follow all in-house safety procedures for working with radiolabeled materials.
- C. All isotopes and work stations where isotopes are used should be controlled access areas. Any one with access to the area must also receive radiation safety training.
- D. Freezers where the isotopes are stored must be locked.
- E. Personal protective equipment must include lab coats (designated specifically for use with radioactive materials), safety glasses, and gloves.
- F. Radioactive materials will only be handled and manipulated in designated areas, which have been clearly identified and labeled accordingly.
- G. Work with source radiation material must be conducted in a fume hood.
- H. Radioactive materials will be stored and/or carried in secondary containment.
- I. When possible, disposable supplies such as pipet tips, absorbent paper, and kim wipes will be used so that contaminated supplies can be readily disposed of as radioactive waste.
- J. Wipe surveys will be conducted at the end of each experiment as well as monthly to ensure that there is no contamination in the laboratory.
- K. The filter plates used in the assay will be designated as solid radioactive waste, while the washes from the filter plates (containing buffer and unbound ^3H -STX) will be handled as liquid radioactive waste. There will be a dry active waste container to hold contaminated items such as the plates, gloves, absorbent paper and kim wipes. There will be a liquid waste jug to hold the contaminated liquid radioactive waste.
- L. All wastes must be disposed of according to state and local laws.

Appendix I. Abbreviations and Acronyms

³ H-STX	Tritiated saxitoxin
AOAC	Association of Analytical Communities
ARC	American Radiolabeled Chemicals
B	Bound CPM
B ₀	Maximum bound CPM
CFSAN	Center for Food Safety & Applied Nutrition
CPM	Counts per minute
diHCl	Dihydrochloride
Eq	Equivalents
HCl	Hydrochloric acid
IC ₅₀	Inhibitory concentration at which CPMs are at 50% max
LC-FD	Liquid chromatography with fluorescence detection
LOD	Limit of detection
LOQ	Limit of quantitation
MBA	Mouse bioassay
MOPS	3-Morpholinopropanesulfonic acid
NaOH	Sodium hydroxide
NIST	National Institute of Standards and Technology
NSSP	National Shellfish Sanitation Program
OMA	Official method of analysis
PMSF	Phenyl methylsulfonyl fluoride
PCOX	Post-column oxidation liquid chromatography with fluorescence detection
Pre-COX	Pre-column oxidation liquid chromatography with fluorescence detection
PSP	Paralytic shellfish poisoning
PSTs	Paralytic shellfish toxins
QC	Quality control
QS	Quality System
RBA	Receptor binding assay
RSD	Relative standard deviation
SLV	Single laboratory validation
STX	Saxitoxin

Single-Laboratory Validation of the Microplate Receptor Binding Assay for Paralytic Shellfish Toxins in Shellfish

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A single-laboratory validation (SLV) study was conducted for the microplate receptor binding assay (RBA) for paralytic shellfish poisoning (PSP) toxins in shellfish. The basis of the assay is the competition between [^3H]saxitoxin (STX) and STX in a standard or sample for binding to the voltage dependent sodium channel. A calibration curve is generated by the addition of 0.01–1000 nM STX, which results in the concentration dependent decrease in [^3H]STX-receptor complexes formed and serves to quantify STX in unknown samples. This study established the LOQ, linearity, recovery, accuracy, and precision of the assay for determining PSP toxicity in shellfish extracts, as performed by a single analyst on multiple days. The standard curve obtained on 5 independent days resulted in a half-maximal inhibition (IC_{50}) of 2.3 nM STX \pm 0.3 (RSD = 10.8%) with a slope of 0.96 \pm 0.06 (RSD = 6.3%) and a dynamic range of 1.2–10.0 nM. The LOQ was 5.3 μg STX equivalents/100 g shellfish. Linearity, established by quantification of three levels of purified STX (1.5, 3, and 6 nM), yielded an r^2 of 0.97. Recovery from mussels spiked with three levels (40, 80, and 120 μg STX/100 g) averaged 121%. Repeatability (RSD_r), determined on six naturally contaminated shellfish samples on 5 independent days, was 17.7%. A method comparison with the AOAC mouse bioassay yielded $r^2 = 0.98$ (slope = 1.29) in the SLV study. The effects of the extraction method on RBA-based toxicity values were assessed on shellfish extracted for PSP toxins using the AOAC mouse bioassay method (0.1 M HCl) compared to that for the precolumn oxidation HPLC method (0.1% acetic acid). The two extraction methods showed linear correlation ($r^2 = 0.99$), with the HCl extraction method yielding slightly higher toxicity values (slope = 1.23). A similar relationship was

observed between HPLC quantification of the HCl- and acetic acid-extracted samples ($r^2 = 0.98$, slope 1.19). The RBA also had excellent linear correlation with HPLC analyses ($r^2 = 0.98$ for HCl, $r^2 = 0.99$ for acetic acid), but gave somewhat higher values than HPLC using either extraction method (slope = 1.39 for HCl extracts, slope = 1.32 for acetic acid). Overall, the excellent linear correlations with the both mouse bioassay and HPLC method and sufficient interassay repeatability suggest that the RBA can be effective as a high throughput screen for estimating PSP toxicity in shellfish.

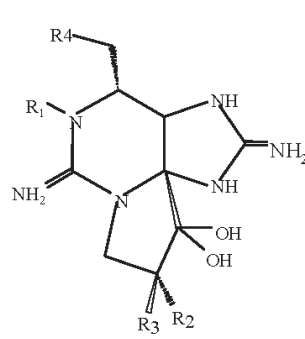
Paralytic shellfish poisoning (PSP) is a seafood intoxication caused by the consumption of shellfish tainted with saxitoxins (STXs) produced by certain species of harmful algae. Saxitoxins are a suite of heterocyclic guanidinium toxins, of which currently more than 21 congeners are known (Figure 1). These congeners occur in varying proportions in the dinoflagellates that produce them and are further metabolized in shellfish that accumulate them, making analytical determination of PSP toxins in shellfish complex. The long-standing regulatory method for PSP toxins is the AOAC mouse bioassay (1), with a regulatory limit of 80 $\mu\text{g}/100$ g shellfish generally applied. Increasing resistance to whole animal testing has driven the need to develop alternative methods suitable for use in a high throughput monitoring or regulatory setting. In the past decade, several alternatives to the mouse bioassay have been developed and validated to various degrees. The precolumn oxidation HPLC method (2) has received First Action approval by AOAC as an Official Method for PSP (2005.06; 3) and has been accepted into the European Food Hygiene Regulations as an alternative to the mouse bioassay and further refined to optimize its use in the United Kingdom Official Control monitoring of PSP toxins in mussels (4). However, although the HPLC method performs well quantitatively, it is quite time consuming for high throughput screening needed by many monitoring programs. A qualitative lateral flow antibody test for PSP toxins with a detection limit of 40 $\mu\text{g}/100$ g, developed by

Jellett Rapid Testing Ltd (Chester Basin, NS, Canada), has been approved in the United States by the Interstate Shellfish Sanitation Conference and the U.S. Food and Drug Administration (FDA) as a screening method. This method performed well in a comparison study with the mouse bioassay, with a false-positive rate of 6% and a false-negative rate of <0.1% (5), but it has not been put through a full AOAC collaborative trial, and does not provide quantitative analysis. To date, a suitable quantitative, high throughput alternative to the mouse bioassay has not been validated through the AOAC Official Methods Program. The current study establishes the single laboratory performance characteristics of the microplate receptor binding assay (RBA) for PSP toxins in shellfish and identifies it as a candidate for fulfilling the requirements of high throughput, quantitative analysis that measures a composite toxic potency in a manner analogous to the mouse bioassay.

STX elicit their paralytic effects by binding to site 1 on the voltage dependent sodium channel, thereby blocking the transmission of neuronal and muscular action potentials. Because all STX congeners bind to site 1 with affinities proportional to their mouse intraperitoneal (IP) toxicity (6), a receptor binding competition assay can be used to measure the integrated toxic potency of STX congeners in a sample, independent of which toxin congeners are present. Moreover, any toxin metabolites originating in the shellfish matrix will also be detected by the assay according to their affinity for the sodium channel receptor. In this binding competition assay, [³H]STX competes with unlabeled STX and/or its derivatives for a finite number of available receptor sites in a rat brain membrane preparation. Following establishment of binding equilibrium, unbound [³H]STX is removed by filtration and

bound [³H]STX is quantified by liquid scintillation counting. The percent reduction in [³H]STX binding in the presence of unlabeled toxin is directly proportional to the amount of unlabeled toxin present. A standard curve is established using increasing concentrations of unlabeled STX, and the concentration of PSP toxins in an unknown sample is quantified using this standard curve.

The assay tested in this single laboratory trial is a modification of the method of Doucette et al. (7) to a 96-well microplate format described by Van Dolah et al. (8). Application of microplate scintillation counting to the PSP assay was first reported by Powell and Doucette (9), who applied it to phytoplankton analysis. The use of the microplate format, in conjunction with microplate scintillation counting, makes the assay suitable for use in a high throughput monitoring or regulatory setting. Several versions of the PSP receptor binding assay have undergone method comparisons in different laboratories with favorable correlations to the mouse bioassay and/or other assays for PSP toxins in shellfish. Suarez-Isla and Valez (10) showed excellent linear correlation ($r^2 = 0.97$) between the RBA and mouse bioassay of 41 shellfish extracts between 40 and 10 000 μg STX equivalents/100 g. Llewellyn et al. (11) found that the sodium channel receptor assay compared well to three other methods of analysis for PSP toxins in shellfish (HPLC, mouse bioassay, and N2A cytotoxicity assay). Ruberu et al. (12) optimized the microplate format assay for use in the Packard Top Count microplate scintillation counter (a single channel counter; GMI, Inc., Ramsey, MN), compared results with the same assay performed on the Wallac microplate counter (a two-channel coincidence counter; Perkin Elmer Wallace, Gaithersburg, MD), and provided further correlation data with



		R1	R2	R3	R4	MU/ μmol
Carbamate	STX	H	H	H	OCONH ₂	2483
	Neo STX	OH	H	H	OCONH ₂	2295
	GTX1	OH	OSO ₃ -	H	OCONH ₂	2468
	GTX2	H	OSO ₃ -	H	OCONH ₂	892
	GTX3	H	H	OSO ₃ -	OCONH ₂	1584
	GTX4	OH	H	OSO ₃ -	OCONH ₂	1803
Sulfocarbamoyl	GTX5 (B1)	H	H	H	OCONHSO ₃ -	160
	GTX6 (B2)	OH	H	H	OCONHSO ₃ -	-
	C1	H	OSO ₃ -	H	OCONHSO ₃ -	15
	C2	H	H	OSO ₃ -	OCONHSO ₃ -	239
	C3	OH	OSO ₃ -	H	OCONHSO ₃ -	33
C4	OH	H	OSO ₃ -	OCONHSO ₃ -	143	
Decarbamoyl	dcSTX	H	H	H	OH	1274
	dcNeoSTX	OH	H	H	OH	-
	dcGTX1	OH	OSO ₃ -	H	OH	-
	dcGTX2	H	OSO ₃ -	H	OH	1617
	dcGTX3	H	H	OSO ₃ -	OH	1872
dcGTX4	OH	H	OSO ₃ -	OH	-	
Deoxydecarbamoyl	doSTX	H	H	H	H	-
	doGTX2	H	H	OSO ₃ -	H	-
	doGTX3	H	OSO ₃ -	H	H	-

Figure 1. Structures and toxic potency of 21 saxitoxin congeners. Toxic potency is listed as mouse units (MU)/ μmole , where a mouse unit is defined as the minimum amount required to kill a 20 g mouse in 15 min when administered by IP injection. The table is modified from ref. 15.

	1	2	3	4	5	6	7	8	9	10	11	12
A	10 ⁻⁶	10 ⁻⁶	10 ⁻⁶	REF	REF	REF	U3 1:10	U3 1:10	U3 1:10	U5 1:200	U5 1:200	U5 1:200
B	10 ⁻⁷	10 ⁻⁷	10 ⁻⁷	QC	QC	QC	U3 1:50	U3 1:50	U3 1:50	U6 1:10	U6 1:10	U6 1:10
C	3 x 10 ⁻⁸	3 x 10 ⁻⁸	3 x 10 ⁻⁸	U1 1:10	U1 1:10	U1 1:10	U3 1:200	U3 1:200	U3 1:200	U6 1:50	U6 1:50	U6 1:50
D	10 ⁻⁸	10 ⁻⁸	10 ⁻⁸	U1 1:50	U1 1:50	U1 1:50	U4 1:10	U4 1:10	U4 1:10	U6 1:200	U6 1:200	U6 1:200
E	3 x 10 ⁻⁹	3 x 10 ⁻⁹	3 x 10 ⁻⁹	U1 1:200	U1 1:200	U1 1:200	U4 1:50	U4 1:50	U4 1:50	U7 1:10	U7 1:10	U7 1:10
F	10 ⁻⁹	10 ⁻⁹	10 ⁻⁹	U2 1:10	U2 1:10	U2 1:10	U4 1:200	U4 1:200	U 1:200	U7 1:50	U7 1:50	U7 1:50
G	10 ⁻¹⁰	10 ⁻¹⁰	10 ⁻¹⁰	U2 1:50	U2 1:50	U2 1:50	U5 1:10	U5 1:10	U5 1:10	U7 1:200	U7 1:200	U7 1:200
H	10 ⁻¹¹	10 ⁻¹¹	10 ⁻¹¹	U2 1:200	U2 1:200	U2 1:200	U5 1:50	U5 1:50	U5 1:50			

U = unknown sample

Figure 2. Standardized plate layout recommended for the microplate RBA for PSP toxins in shellfish extracts. U = unknown sample.

the mouse bioassay. Usup et al. (13) utilized the microplate RBA method to compare predicted toxicity values in samples spiked with different STX congeners as assayed by the mouse bioassay and the RBA. Llewellyn (14) defined the competitive behavior of PSP toxin mixtures in receptor binding assays, using both the sodium channel and saxiphilin receptors, which explains their composite toxicity. However, none of these previous studies fully characterized assay performance according to AOAC single-laboratory validation (SLV) criteria that are the underpinning required for proceeding with an AOAC collaborative trial. Therefore, the current study was carried out to fulfill those requirements.

Experimental

Apparatus

- (a) *Microplate scintillation counter*.—Wallac Microbeta, GMI Inc. (Ramsey, MN).
- (b) *Microplate filtration manifold*.—Millipore (Bedford, MA).
- (c) *Hot plate*.—Fisher Scientific (Suwanee, GA).
- (d) *Countertop centrifuge*.—For 15 mL tubes, capable of 3000 × g (Fisher Scientific).
- (e) *Microtiter filter plates (96 well) with 1.0 μm pore size type FB glass fiber filter/0.65 μm pore size Duropore support membrane*.—Cat. No. MSFB N6B 50 (Millipore Corp., Billerica, MA).
- (f) *Microplate sealing tape*.—Cat. No. MATA HCL00 (Millipore Corp.).

(g) *Vortex mixer*.—Daigger Vortex Genie II (Daigger Scientific, Vernon Hills, IL).

(h) *Teflon/glass tissue homogenizer*.—Wheaton (Millville, NJ).

(i) *Polytron homogenizer*.—Brinkmann Instruments (Westbury, NY).

Reagents

- (a) *Hydrochloric acid (HCl)*.—0.1 M.
- (b) [³H]STX.—0.1 mCi/mL, ≥10 Ci/mmol, ≥90% radiochemical purity (International Isotopes Clearinghouse, Leawood, KS).
- (c) *STX diHCl*.—FDA reference standard (Office of Seafood, Laurel, MD) or National Research Council (NRC) of Canada Institute of Marine Biosciences (Halifax, NS, Canada).
- (d) *Assay buffer*.—75 mM HEPES [4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid; Cat. No. H9136]/140 mM NaCl, pH 7.5 (Sigma, St. Louis, MO).
- (e) *Liquid scintillation cocktail*.—Optiphase (PerkinElmer Life Sciences, Downers Grove, IL).

Preparation of Samples (0.1 M HCl Extraction)

Shellfish samples were shucked and homogenized according to the AOAC mouse bioassay protocol (1). For the HCl extraction method, 5.0 (±0.1) g of tissue homogenate was transferred to a tared 15 mL conical polypropylene centrifuge tube. A 5.0 mL volume of 0.1 M HCl was added, and the sample was mixed on a Vortex mixer. The pH was checked to

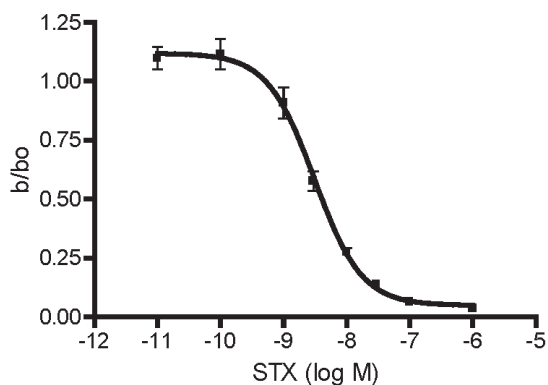


Figure 3. Average of five calibration curves obtained by one analyst in five independent assays on separate days. $IC_{50} = 2.23 \pm 0.23$ nM, slope = 0.96 ± 0.06 , error bars are \pm SD.

confirm it was between 3.0 and 4.0 in order to avoid alkalization and destruction of the toxin, and adjusted with 1 M HCl or 0.1 M NaOH as needed. Tubes were placed in a beaker of boiling water on a hot plate for 5 min with the caps loosened. Following removal from the boiling water bath, samples were allowed to cool to room temperature, and the pH was again confirmed to be between 3.0 and 4.0. The entire contents were then transferred to a graduated cylinder, diluted volumetrically to 10 mL, and centrifuged for 5 min at $1000 \times g$. The supernatant was transferred to a clean tube.

Preparation of Samples (Acetic Acid Extraction Method)

In a 50 mL plastic centrifuge tube, 5.0 ± 0.1 g homogenate was mixed with 3.0 mL 1% acetic acid on a vortex mixer. Tubes were capped loosely to avoid pressure buildup and placed in a boiling water bath for 5 min. Following removal from the water bath, samples were cooled in a beaker of cold water for 5 min, and then centrifuged for 10 min at $3000 \times g$. The supernatant was transferred to a 15 mL graduated conical test tube. A 3 mL amount of 1% acetic acid was added to the original tube with solid residue, mixed well on a vortex mixer, and centrifuged again for 10 min at $3000 \times g$. The second supernatant was combined with the first and diluted to 10 mL with water.

Preparation of Stock Solutions, Standards, and Reagents for Assay

(a) *Radioligand solution.*— $[^3H]$ STX stock is provided in 50 μ Ci ampules, 24 Ci/mmol, 0.1 mCi/mL (4.17 μ M). A 15 nM working stock of $[^3H]$ STX was prepared fresh daily in 75 mM HEPES/140 mM NaCl (for 2.5 nM final in-well concentration).

(b) *STX standard curve.*—FDA STX dihydrochloride reference standard (100 μ g/mL or 268.8 μ M) used to prepare a bulk standard curve made up in advance and stored at 4°C for up to 1 month. The stock standard curve was made consisted of eight concentrations of STX in 0.003 M HCl [6×10^{-6} , 6×10^{-7} , 1.8×10^{-7} , 6×10^{-8} , 1.8×10^{-8} , 6×10^{-9} , 6×10^{-10} , $6 \times$

Table 1. RBA measurements of calibration standards for assay linearity assessment (nM STX; $n = 5$)

Nominal	Mean	SD	RSD
1.5	1.7	0.16	10
3.0	3.0	0.52	17
6.0	6.0	0.34	6

10^{-11} , and 0.003 M only HCl (reference)], which when diluted 1:6 in the assay, resulted in a standard curve of 0.01 nM–1000 nM STX. The reference provided a measure of total $[^3H]$ STX binding in the absence of unlabeled STX.

(c) *Calibration standard (QC check).*—A reference standard containing 1.8×10^{-8} M STX standard (3.0×10^{-9} M STX in assay) was prepared in 0.003 M hydrochloric acid, aliquotted in 1 mL volumes, and stored at 4°C for routine use (stable up to 1 month). On the day of the assay, 200 μ L of each standard were pipetted into mini-dilution tubes for ease of pipetting into the microplate using an eight-channel pipettor.

(d) *Rat brain membrane homogenate.*—Cerebral cortices from 6-week-old male Holzman rats (Harlan Bioproducts, Indianapolis, IN) were homogenized on ice in a glass/Teflon tissue homogenizer in 75 mM HEPES/140 mM NaCl, pH 7.5, containing 0.1 mM PMSF (phenylmethanesulfonylfluoride; 12.5 mL/brain) at 385 rpm for 10 strokes. Pooled homogenates were centrifuged at $20\,000 \times g$ for 15 min at 4°C and the pellet was resuspended in HEPES buffer (12.5 mL/brain) and rehomogenized on ice using a Polytron homogenizer set at 70% power for 20 s to ensure a fine suspension. The brain homogenate was aliquotted 2 mL/tube in cryovials and stored at -80°C . The protein concentration of the brain homogenate was determined using the Micro bicinchoninic acid (BCA) Assay (Pierce, Rockford, IL). For each assay, an aliquot of brain homogenate was thawed on ice and diluted with ice cold 75 mM HEPES/150 mM NaCl, pH 7.5, to yield a final protein concentration of 0.5 mg/mL in the assay.

Table 2. Recovery of analyte from spiked samples (μ g STX equiv./100 g)

Nominal	Mean	SD	Measured RSD _r	Recovery, %
0	<dl ^a			
40	47	8.6	18.7	115
80	103.7	21.8	21	129
120	145.5	15.2	10.5	121

^a <dl = Less than LOQ (5 μ g STX equiv./100 g).

Table 3. Comparison of receptor binding assay (RBA; n = 5) with AOAC mouse bioassay (MBA) of naturally contaminated shellfish (µg STX equiv./100 g)

Sample	MBA	RBA mean	SD	RSD
LP1	340	438	74	17
LP2	534	715	96	13
LP3	1158	1533	329	21
LP4	65	91	7	9
LP5	350	608	150	25
LP6	462	518	114	22

Assay Procedure

(a) *Plate setup and incubation.*—A standardized plate layout was used for all assays (Figure 2). All standards, reference, QC check, and shellfish extracts were run in triplicate wells. For shellfish extracts, a standardized dilution series was run for each sample (1:10, 1:50, and 1:200), which ensured that at least one dilution would fall on the linear part of the competition curve for shellfish that contains between approximately 5 and 1500 µg STX equiv./100 g. Reagents were added in the following order: 35 µL STX standard or sample, then 35 µL [³H]STX, followed by 140 µL brain homogenate. The addition of brain homogenate was carried out with sufficient force to ensure mixing of the well contents, but without risk of splashing. The plate was then covered and incubated at 4°C for 1 h.

(b) *Assay filtration and counting.*—The plate was filtered using a microplate vacuum filtration manifold, and each well rinsed twice with 200 µL ice-cold HEPES buffer at a filtration rate that ensured all wells were dry within 2–5 s. The microplate was then placed in a microplate scintillation counter cassette, and the bottom was sealed with plate sealing tape. Lastly, 50 µL scintillation cocktail was added to each well, and the top of the plate was sealed with sealing tape. The plate was allowed to sit for 30 min to ensure impregnation of the filters with scintillant prior to counting for 1 min/well in the microplate scintillation counter.

Data Analysis

Curve fitting was performed using a four-parameter logistic curve fitting model for a one-site receptor binding using Wallac Multicalc software. The software reports the in-well sample concentration in nM equiv. STX. Sample concentration was then calculated in µg STX equivalents/100 g shellfish using the following formulas:

$$\begin{aligned}
 & (\text{nM equiv. STX}) \times (\text{sample dilution}) \times \frac{(210 \mu\text{L total volume})}{35 \mu\text{L sample}} \\
 & = \text{nM equiv. STX in extract}
 \end{aligned}$$

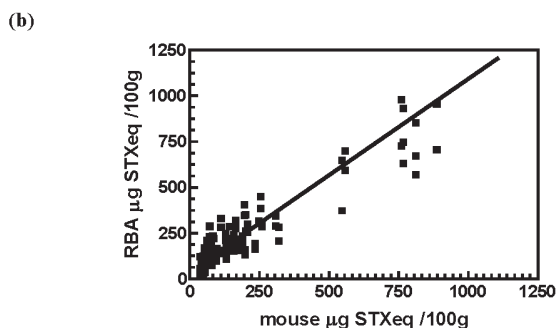
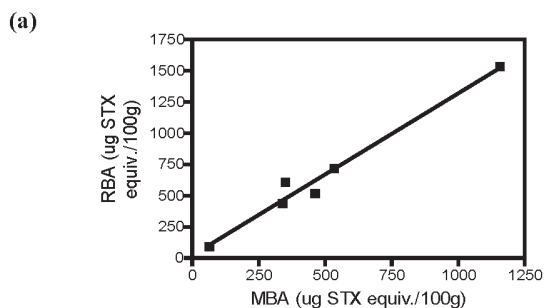


Figure 4. Linear correlation analysis between the RBA and mouse bioassay. (a) Average values of six naturally contaminated samples analyzed on five independent RBA assay days ($r^2 = 0.98$, slope = 1.29). (b) A separate study of 110 shellfish extracts analyzed by RBA and MBA yielded an r^2 of 0.88 with a slope of 1.32.

$$\begin{aligned}
 & (\text{nm equiv. STX in extract}) \times \frac{1 \text{ L}}{1000 \text{ mL}} \times \frac{372 \text{ ng}}{\text{nmol}} \times \frac{1 \mu\text{g}}{1000 \text{ ng}} \\
 & = \mu\text{g STX equiv./mL} \\
 & \mu\text{g STX equiv./mL} \times \frac{\text{mL extract}}{\text{g shellfish extracted}} \times 100 \\
 & = \mu\text{g STX equiv./100 g shellfish}
 \end{aligned}$$

Critical Control Points

(1) For a ligand that interacts specifically at one receptor site, the slope of the resulting competition curve should theoretically be 1.0. If the slope of the curve for a given assay is outside of the acceptable range of 0.8–1.2, linearity of the assay will be compromised, and quantification of the unknowns will be incorrect. Therefore, the assay should be re-run.

(2) The QC check standard should fall within ±30% of the stated value (3.0 nM). If the QC check standard does not fall within acceptable limits, the assay should be re-run.

Table 4. RBA-determined toxicities of nine naturally contaminated shellfish homogenates extracted using the 0.1 M HCl extraction method or the 1% acetic acid extraction method (μg STX equiv./100 g)

Sample	HCl			Acetic acid		
	Mean	SD	RSD	Mean	SD	RSD
1	11	4	36	19	7	39
2	600	143	24	488	104	21
3	690	142	21	584	167	29
4	136	8	6	131	41	31
5	152	27	18	167	21	13
6	302	87	29	270	72	27
7	340	88	26	264	63	24
8	262	79	30	252	48	19
9	63	26	41	54	19	34

(3) Sample quantification should be done only on dilutions that on the linear part of the curve [$b/b_0 = 0.2-0.7$, where B is the bound counts/min (CPM) in the sample and B_0 is the maximum CPM]. The RSD of the CPM must be $<30\%$.

(4) For a given sample, if none of the sample dilutions falls within the linear range (i.e., the concentration is too high, $b/b_0 < 0.2$), further dilutions must be made and the sample reanalyzed if a quantitative value is desired. If the sample concentration is too low to be quantified (i.e., $b/b_0 > 0.7$) at sample dilution 1:10, the sample must be reported as below the LOQ.

Mouse Bioassay and HPLC Procedures

Shellfish samples extracted in parallel using the HCl and acetic acid extraction methods described above were analyzed using the standard protocols prescribed by the AOAC methods for mouse bioassay (1) or precolumn oxidation HPLC method (2).

Results and Discussion

Calibration Curve

To establish the dynamic range and repeatability of the calibration curve, five assays were performed by one analyst on separate days. The composite curve (Figure 3) resulted in a half-maximal inhibition (IC_{50}) of $2.3 \text{ nM STX} \pm 0.3$ (RSD = 10.8%) with a slope of 0.96 ± 0.06 (RSD = 6.3%). Using the linear part of the curve ($0.2-0.7 b/b_0$) for quantification, a dynamic range of approximately one order of magnitude, $1.2-10.0 \text{ nM STX}$, was observed, as expected for a one-site binding assay. A QC check sample (3.0 nM STX) run in each assay averaged $3.0 \pm 0.5 \text{ nM}$ (RSD_r = 17.3%), with a recovery of 99.3% .

LOQ

Shellfish extracts were diluted a minimum of 10-fold prior to analysis to minimize matrix effects that can result in false positives. The LOQ was empirically determined as the

concentration, in a 10-fold diluted sample, that results in a b/b_0 of 0.7. This is a more conservative cutoff than the $0.8 b/b_0$ frequently used in receptor assays and was used because quantification was unacceptably variable above this b/b_0 cutoff. This results in an LOQ of approximately $5 \mu\text{g}$ equiv. STX/100 g shellfish, which provides a more than one order of magnitude margin relative to the regulatory limit of $80 \mu\text{g}/100 \text{ g}$.

Linearity

Linearity was assessed by five independent assays of three calibration standards that were expected to fall on the curve between 0.2 and $0.7 b/b_0$: $1.5, 3.0,$ and 6.0 nM STX prepared from FDA STX diHCl standard. Expected and measured values are listed in Table 1. Linear regression yielded a slope of 0.98 and an r^2 of 0.97 .

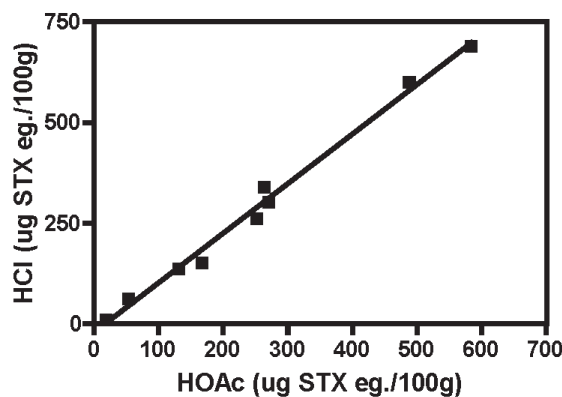


Figure 5. Linear correlation between HCl and acetic acid (HOAc) extracts analyzed by RBA. Results are average values of nine naturally contaminated samples obtained from four independent assays ($r^2 = 0.99$, slope = 1.23).

Table 5. HPLC analysis of nine naturally contaminated samples (1–9) extracted using 0.1 M HCl^a

Sample	STX	NEO ^b	GTX1,4 ^c	GTX2,3	B1	C1,2	Total PSP	As STX equivalent
HCl-1	3.5	0.0	0.0	7.3	0.0	0.0	10.8	6
HCl-2	231.6	23.9	42.6	324.5	28.3	249.7	900.4	412
HCl-3	220.8	53.7	74.9	436.3	43.4	338.1	1167.2	494
HCl-4	48.3	2.7	8.6	85.1	10.7	17.1	172.5	90
HCl-5	86.5	1.1	0.0	64.7	14.9	11.3	178.5	113
HCl-6	114.5	0.0	0.0	166.6	15.1	36.8	333.0	180
HCl-7	96.4	10.1	72.9	398.7	9.3	36.1	623.5	304
HCl-8	84.6	6.0	32.8	225.7	4.9	18.5	372.5	197
HCl-9	11.2	0.0	6.1	47.9	0.0	0.0	65.2	33

^a Values are in $\mu\text{g}/100\text{ g}$, as specific PSP congener or its STX equivalents, as indicated by the column headers.

^b NEO = Neosaxitoxin.

^c GTX = Gonyautoxin.

Recovery

Mussel tissue homogenates obtained from a local market were spiked with FDA STX diHCl standard at four levels bracketing the regulatory limit (0, 40, 80, and 120 $\mu\text{g}/100\text{ g}$) followed by thorough homogenization using a Polytron blender. Aliquots of spiked homogenate were stored at -80°C until extraction in 0.1 M HCl according to the protocol in the *Experimental* section. Extracts were analyzed in five assays performed on independent days. The mean recovery was 121% (Table 2).

Comparison of RBA-Reported Toxicity with the AOAC Mouse Bioassay

Six naturally contaminated shellfish samples were extracted in 0.1 M HCl according to the protocol in the *Experimental* section, and analyzed in five assays on

independent days (Table 3). Three shellfish species were represented: clam *Mya arenaria* (whole) LP1, LP4; mussel *Mytilus edulis* (whole) LP2, LP3; and scallop *Plactopecten magellanicus* (viscera) LP5, LP6. Between-assay RSDs ranged from 9 to 25% (mean 17.7%). An r^2 of 0.98 was obtained relative to the mouse bioassay, with a slope of 1.29 (Figure 4a).

A separate study of 110 naturally contaminated shellfish samples, extracted using the 0.1 M HCl method, and analyzed by RBA and mouse bioassay, yielded similar results with an r^2 of 0.88 and a slope of 1.32 (Figure 4b).

Effect of Extraction Method on RBA-Reported Toxicities

The recent approval of the precolumn oxidation HPLC method for PSP toxins as AOAC Official Method **2005.06** (3) and its potential recognition as a reference method for PSP

Table 6. HPLC analysis of the same nine naturally contaminated samples (1–9) extracted using 1% acetic acid^a

Sample	STX	NEO	GTX1,4	GTX2,3	B1	C1,2	Total PSP	As STX equivalent
HOAc-1	3.4	0.0	0.0	7.3	0.0	0.0	10.7	6
HOAc-2	187.6	13.1	21.7	280.7	25.1	248.9	777.1	329
HOAc-3	175.2	35.6	79.2	335.9	37.2	237.7	900.9	393
HOAc-4	33.4	3.1	11.3	61.8	6.0	15.5	131.1	68
HOAc-5	59.3	3.1	0.0	67.6	10.8	19.3	160.0	89
HOAc-6	100.8	0.0	0.0	158.0	11.8	28.4	299.0	162
HOAc-7	67.4	11.2	42.7	228.4	5.2	15.6	370.5	192
HOAc-8	71.0	8.3	34.4	190.3	4.3	12.6	320.8	173
HOAc-9	11.2	0.0	11.7	38.1	0.0	61.0	122.1	33

^a Values are in $\mu\text{g}/100\text{ g}$, as specific PSP congener or its STX equivalents, as indicated by the column headers.

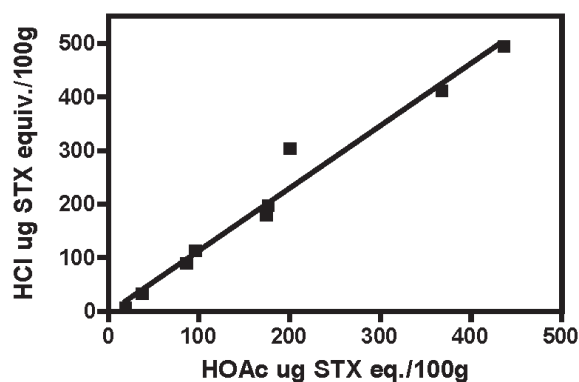


Figure 6. Linear correlation between HCl and acetic acid (HOAc) extracts analyzed by HPLC (slope = 1.16, $r^2 = 0.97$).

toxins prompted an investigation of the effects of extraction method on toxicity values reported by the RBA. Whereas the AOAC mouse bioassay prescribes shellfish extraction in 0.1 M HCl, the HPLC method uses extraction in 1% acetic acid. The 0.1 M HCl extraction procedure is known to result in the partial conversion of certain low-toxicity sulfocarbamoyl congeners to more highly toxic congeners in shellfish extracts, especially gonyautoxins, GTX5 and GTX6, to STX and neoSTX, and, thus, may result in somewhat higher toxicity values. To assess the effects of extraction procedure on RBA-reported toxicity, nine naturally contaminated shellfish samples (six blue mussel and three scallop) were homogenized and extracted independently using 0.1 M HCl and 1% acetic acid as described in the *Experimental* section. PSP toxicity in the extracts was then determined in four RBA assays run on independent days (Table 4). The between-assay RSD did not differ for samples prepared using the two extraction methods (25.8 and 26.3%, respectively). In general, the HCl extraction method resulted in slightly higher total toxicity values than reported for the acetic acid extracts (slope 1.23, $r^2 = 0.99$; Figure 5). The higher values reported for the HCl extracts are not explained by the conversion of sulfocarbamoyl toxins to more potent congeners in the HCl extracts, as can be seen in the toxin profiles determined by HPLC (Tables 5 and 6). Rather, the recovery of most congeners appears to be higher in the HCl extract. The higher concentrations reported in the HCl extract may reflect differences in the method by which volume is adjusted in the two extraction procedures. In the HCl method, final extract volume adjustment is made with the shellfish matrix present. In the acetic acid extraction, the matrix is first removed, the pellet re-extracted, the two extracts pooled, and then the final volume adjusted. HPLC analysis of the same samples showed a similar relationship between values reported for the HCl and acetic acid extracts (slope = 1.16, $r^2 = 0.97$; Figure 6) as seen in the RBA, with the HCl extracts containing greater STX equivalent/100 g.

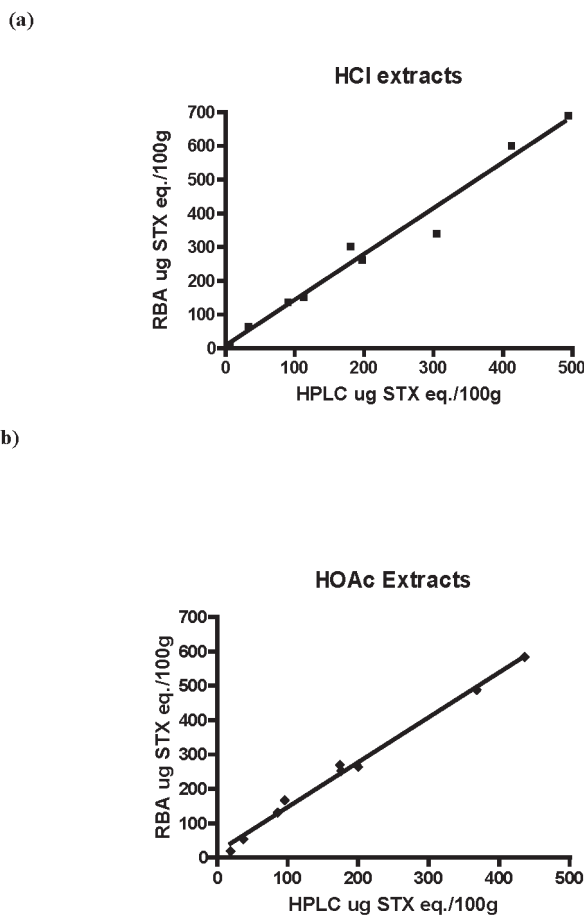


Figure 7. Linear correlation between RBA and HPLC for samples extracted (a) by the HCl method ($r^2 = 0.98$, slope = 1.39) and (b) by the acetic acid method ($r^2 = 0.99$, slope = 1.32).

Comparison of RBA with HPLC

The RBA showed good linear correlation with HPLC analysis of both HCl ($r^2 = 0.98$, slope = 1.39) and acetic acid ($r^2 = 0.99$, slope = 1.32) extracts, in both cases giving somewhat higher toxicities than the HPLC method (Figure 7). A number of factors may contribute to the difference in results for total toxic potencies by these two methods. The higher toxicity values given by the RBA may result in part from the fact that the HPLC method uses the STX free base molecular weight (300 Da), whereas the receptor assay (and mouse bioassay) uses the STX dihydrochloride molecular weight (372 Da) to calculate concentration, which would result in approximately 20% higher values in the RBA. Additional differences may result from the use of FDA as compared to the NRC saxitoxin standards in the RBA and HPLC methods, respectively. Higher RBA results may also result from the dominance of the more potent PSP congeners over the weaker congeners in mixtures competing for binding to the receptor, as detailed in ref. 13, which reflects their binding affinities. In

contrast to this complex behavior, the HPLC method adds linearly the concentrations of each congener based on toxic potencies determined by mouse bioassay for isolated congeners. In some cases, e.g., 11-hydroxysulfate epimers, the concentrations of separate epimers pairs are not resolved by HPLC, although their potencies differ widely as do their ratios in shellfish samples. Lastly, higher toxicity values reported by the RBA may reflect the presence of congeners or metabolites not reported by the HPLC method.

Ruggedness

Although formal ruggedness testing was not carried out during this SLV study, several steps in the procedure might be noted that can affect the precision and accuracy of the results. First, it is important to clarify shellfish extracts by centrifugation prior to running the assay, particularly if extracts are stored refrigerated or frozen before analysis, as precipitates in the extract may cause nonspecific binding that may result in overestimates of PSP toxin concentrations. Second, since the rat brain homogenate is a suspension, it is important to ensure that it remains evenly suspended by frequent vortex mixing or pipetting prior to and during its addition to the plate. The rate of assay plate filtration should ensure that the wells clear in 2–5 s, and the rinse buffer should be ice cold in order to minimize the rate of toxin release from the receptor. Lastly, following addition of liquid scintillant to the microplate wells, it is essential to allow a minimum of 30 min for the scintillant to penetrate the filters before counting. Counting prematurely can result in increased variability between wells and lower counts/well, thus increasing RSD. A count time of 1 min/well was chosen for this study as a compromise between optimum RSD and assay throughput. Increasing the count time to 5 min/well has been shown to improve the between-well RSD in this assay when using the Packard Top Count scintillation counter, a single detector instrument with somewhat lower efficiency than the Wallac Microbeta used in the current study (11).

Summary

This SLV and method comparison study demonstrates excellent linear correlation ($r^2 > 0.98$) between the microplate receptor binding assay and both the mouse bioassay and the precolumn oxidation HPLC method for the determination of PSP toxins in shellfish. The microplate format of the assay, when coupled with microplate scintillation counting, provides a quantitative high throughput screening tool for PSP toxin testing in shellfish. The tendency of the RBA to overestimate PSP toxicity relative to the reference methods minimizes the chance of returning false negatives. Where RBA-measured

toxicity results in STX equivalent values close to the regulatory limit, confirmation with a reference method is necessary if a regulatory decision is being made. Nonetheless, application of the assay as a high throughput screen can alleviate the unnecessarily large numbers of animals used for the mouse bioassay on negative samples and, similarly, alleviate the lengthy analysis of samples by HPLC at very high or very low concentrations. We propose that this method be collaboratively tested to establish if it is robust enough to be used in monitoring and regulatory laboratories.

Acknowledgments

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FOOD CHEMICAL CONTAMINANTS

Appendix III

Determination of Paralytic Shellfish Toxins in Shellfish by Receptor Binding Assay: Collaborative Study

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A collaborative study was conducted on a microplate format receptor binding assay (RBA) for paralytic shellfish toxins (PST). The assay quantifies the composite PST toxicity in shellfish samples based on the ability of sample extracts to compete with ^3H saxitoxin (STX) diHCl for binding to voltage-gated sodium channels in a rat brain membrane preparation. Quantification of binding can be carried out using either a microplate or traditional scintillation counter; both end points were included in this study. Nine laboratories from six countries completed the study. One laboratory analyzed the samples using the precolumn oxidation HPLC method (AOAC Method 2005.06) to determine the STX congener composition. Three laboratories performed the mouse bioassay (AOAC Method 959.08). The study focused on the ability of the assay to measure the PST toxicity of samples below, near, or slightly above the regulatory limit of 800 (μg STX diHCl equiv./kg). A total of 21 shellfish homogenates were extracted in 0.1 M HCl, and the extracts were analyzed by RBA in three assays on separate days. Samples included naturally contaminated shellfish samples of different species collected from several geographic regions, which contained varying STX congener profiles due to their exposure to different PST-producing dinoflagellate species or differences in toxin metabolism: blue mussel (*Mytilus edulis*) from the U.S. east and west coasts, California mussel (*Mytilus californianus*) from the U.S. west coast, chorito mussel (*Mytilus chilensis*) from Chile, green mussel (*Perna canaliculus*) from New Zealand,

Atlantic surf clam (*Spisula solidissima*) from the U.S. east coast, butter clam (*Saxidomus gigantea*) from the west coast of the United States, almeja clam (*Venus antiqua*) from Chile, and Atlantic sea scallop (*Plactopecten magellanicus*) from the U.S. east coast. All samples were provided as whole animal homogenates, except Atlantic sea scallop and green mussel, from which only the hepatopancreas was homogenized. Among the naturally contaminated samples, five were blind duplicates used for calculation of RSD_r . The interlaboratory RSD_R of the assay for 21 samples tested in nine laboratories was 33.1%, yielding a HorRat value of 2.0. Removal of results for one laboratory that reported systematically low values resulted in an average RSD_R of 28.7% and average HorRat value of 1.8. Intralaboratory RSD_r , based on five blind duplicate samples tested in separate assays, was 25.1%. RSD_r obtained by individual laboratories ranged from 11.8 to 34.9%. Laboratories that are routine users of the assay performed better than nonroutine users, with an average RSD_r of 17.1%. Recovery of STX from spiked shellfish homogenates was 88.1–93.3%. Correlation with the mouse bioassay yielded a slope of 1.64 and correlation coefficient (r^2) of 0.84, while correlation with the precolumn oxidation HPLC method yielded a slope of 1.20 and an r^2 of 0.92. When samples were sorted according to increasing toxin concentration (μg STX diHCl equiv./kg) as assessed by the mouse bioassay, the RBA returned no false negatives relative to the 800 μg STX diHCl equiv./kg regulatory limit for shellfish. Currently, no validated methods other than the mouse bioassay directly measure a composite toxic potency for PST in shellfish. The results of this interlaboratory study demonstrate that the RBA is suitable for the routine determination of PST in shellfish in appropriately equipped laboratories.

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The recommendation was approved by the Method Committee on Paralytic Shellfish Toxins as First Action. See "Methods News," (2011) *Inside Laboratory Management*, January/February issue.

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Paralytic shellfish poisoning (PSP) is caused by a suite of heterocyclic guanidinium toxins collectively called saxitoxins (STXs). Currently more than 21 congeners of STX are known; they occur in varying proportions in the dinoflagellates that produce them and may be further

Table 1. Shellfish homogenate samples analyzed for PSTs in the collaborative study^a

Sample No.	Sample ID	Shellfish species and origin	Blind duplicate
1	MLV05	Atlantic sea scallop (<i>Plactopecten magellanicus</i>) from the U.S. east coast	x
2	MLV06	California mussel (<i>Mytilus californianus</i>) from the U.S. west coast	x
3	MLV08	Green mussel (<i>Perna canaliculus</i>) from New Zealand	
4	MLV09	Blue mussel (<i>M. edulis</i>) from the U.S. west coast	x
5	MLV12	Blue mussel (<i>M. edulis</i>) east coast U.S., spiked with 200 µg/kg STX diHCl	
6	MLV14	Blue mussel (<i>M. edulis</i>) east coast U.S., spiked with 1200 µg/kg STX diHCl	
7	MLV16	Almeja clam (<i>Venus antique</i>) from Chile	
8	MLV01	Surf clam (<i>Spisula solidissima</i>) from the U.S. east coast	
9	MLV02	Chorito mussel (<i>M. chilensis</i>) from Chile	
10	MLV04	Scallop (<i>Plactopecten magellanicus</i>) from the U.S. east coast	
11	MLV07	Blue mussel (<i>M. edulis</i>) east coast U.S.	x
12	MLV09	Blue mussel (<i>M. edulis</i>) from the U.S. west coast	x
13	MLV11	Almeja clam (<i>Venus antique</i>) from Chile clam	x
14	MLV13	Blue mussel (<i>M. edulis</i>) east coast U.S., spiked with 500 µg/kg STX diHCl	
15	MLV03	Chorito mussel (<i>M. chilensis</i>) from Chile	
16	MLV05	Atlantic sea scallop (<i>Plactopecten magellanicus</i>) from the U.S. east coast	x
17	MLV06	California mussel (<i>M. californianus</i>) from the U.S. west coast	x
18	MLV07	Blue mussel (<i>M. edulis</i>) east coast U.S.	x
19	MLV10	Butterclam (<i>Saxidomus gigantea</i>) from the U.S. west coast	
20	MLV11	Almeja clam (<i>Venus antique</i>) from Chile clam	x
21	MLV15	Blue mussel (<i>M. edulis</i>) negative control, east coast U.S.	

^a Sample number identifies the individual samples analyzed in the assays, with 1–7 analyzed in assay 1, 8–14 in assay 2, and 15–21 in assay 3. Sample identification (MLV for multilaboratory validation) describes the 16 unique samples, among which five were assayed as blind duplicates, to make a total of 21 samples. Blind duplicates, run in different assays, are identified by an “x.”

metabolized in shellfish that accumulate them, making analytical determination of paralytic shellfish toxins (PST) in shellfish complex. The long-standing regulatory method for PST is the AOAC mouse bioassay (1; AOAC Method **959.08**), with a regulatory limit of 800 µg STX di HCl equiv./kg shellfish generally applied, but established at 400 µg STX diHCl equiv./kg in certain countries (e.g., the Philippines). However, at concentrations near the regulatory limit, the mouse bioassay can significantly underestimate PST in shellfish (2). This, in addition to increasing resistance to live animal testing in both the United States and the European Union (EU), has increased the need to develop alternative methods suitable for use in a high-throughput monitoring or regulatory setting.

In the past decade, several alternatives to the mouse bioassay have been developed. In the EU, the mouse bioassay remains the reference method for PST in shellfish, but European Commission (EC) Regulation 1664/2006 specifies that other internationally recognized methods may be used. Two HPLC methods, a precolumn oxidation method (3, 4; AOAC Method **2005.06**) and a postcolumn oxidation method (5; AOAC Method **2011.02**), have been approved by AOAC as *Official Methods*SM for PSP toxin analysis. The EC directive recognizes the precolumn oxidation HPLC method (AOAC Method **2005.06**) as an alternative to the mouse bioassay, but retains the mouse bioassay as the reference method in instances where results are challenged. HPLC methods separate and quantify individual

STX congeners, which are then recombined according to their toxic equivalencies to yield a composite PST toxicity value. Although the HPLC methods perform well quantitatively, a high-throughput screening method capable of reporting toxic potency directly is still desirable for monitoring programs that often screen large numbers of negative samples. A qualitative lateral flow antibody test for PST with a reported detection limit of 400 µg STX equiv./kg was developed by Jellett Rapid Testing Ltd (Chester Basin, NS, Canada) and approved by the U.S. Interstate Shellfish Sanitation Conference and the U.S. Food and Drug Administration as a screening method in specific circumstances. This method performed well in a comparison study with the mouse bioassay (6), but is not fully quantitative and has not been subjected to a full AOAC collaborative trial. To date, a suitable quantitative, high-throughput alternative to the mouse bioassay has not been validated through the AOAC Official Methods Program. The receptor binding assay (RBA) for PST is an excellent candidate for fulfilling the requirements of a high-throughput, quantitative assay that directly reports a composite toxic potency.

The basis of the RBA is the interaction between the toxins and their pharmacological target. All STX congeners bind to site 1 on the alpha subunit of the voltage-gated sodium channel with binding affinities proportional to their toxic potency (7). Therefore, an RBA can quantitatively measure the combined toxic potency of mixtures of STX congeners in a sample,

independent of the toxin congeners present (8). In the RBA for PST, tritiated STX ($[^3\text{H}]$ STX) competes with unlabeled STX and/or its congeners for a finite number of available receptor sites in a rat brain membrane preparation. Following establishment of binding equilibrium, unbound $[^3\text{H}]$ STX is removed by filtration and receptor bound $[^3\text{H}]$ STX quantified by liquid scintillation counting. The reduction in $[^3\text{H}]$ STX binding is directly proportional to the amount of unlabeled toxin present. A standard curve is generated using increasing concentrations of nonradiolabeled STX standard from 10^{-10} to 10^{-6} M STX. The concentration of toxin in samples is determined in reference to the standard curve.

The assay being tested in this collaborative trial is a modification of the method of Doucette et al. (9) to incorporate a 96-well microtiter plate format, which increases sample throughput and minimizes error by reducing sample handling and pipetting steps. This microplate PST RBA was evaluated in a single-laboratory validation (SLV) study (10), which established an interassay repeatability (RSD_r) of 17.7% and good correlation with the mouse bioassay and precolumn oxidation HPLC methods. The toxin concentrations in shellfish tested in the SLV study ranged from near to well above the regulatory limit (approximately 900–15 000 μg STX diHCl equiv./kg). The current study focuses more specifically on the performance of the RBA in the critical range of shellfish toxicities below, near, and slightly above the regulatory limit (approximately 150–2400 μg STX diHCl equiv./kg).

The results of the collaborative study suggest that the RBA for PST is a suitable high-throughput screen for PST in shellfish. Although HPLC methods offer quantitative information on congener composition of samples, often the desired information is composite toxic potency, which requires the summation of individual congeners, corrected for their individual toxic equivalencies. The RBA provides a single integrated toxic potency value that reflects activity of all known and potential unknown congeners present in the sample. Use of the microtiter plate format, in conjunction with microplate scintillation counting, provides the ability to screen multiple samples simultaneously in a total assay time of less than 3 h. The assay format described in the current study provides for the quantitative determination of composite PST toxicity in seven shellfish extracts per 96-well microplate, each run in triplicate at three dilutions, covering toxicity ranges of approximately 35–6000 μg STX diHCl equiv./kg. In a high-throughput assay setting, multiple plates can be set up simultaneously, so that six assay plates can easily be accommodated each day by a single analyst, for a throughput of 42 samples/day. This compares favorably to an estimated throughput of 20–25 samples a day by the precolumn HPLC method (B. Niedzwiadek, Health Canada, personal communication) or 30–35 by mouse bioassay (B. Suarez, University of Chile, personal communication).

Collaborative Study

The focus of this study was to assess the performance of the RBA to determine PST toxicity in samples of commercially important shellfish at a range of concentrations below and above the regulatory limit. Twenty-one shellfish homogenates were included in the study, which represented 16 unique samples (Table 1). The homogenates included 12 naturally contaminated shellfish samples of different species collected from several

geographic regions: blue mussel (*M. edulis*) from the U.S. east and west coasts, California mussel (*M. californianus*) from the U.S. west coast, chorito mussel (*M. chilensis*) from Chile, green mussel (*Perna canaliculus*) from New Zealand, Atlantic surf clam (*Spisula solidissima*) from the U.S. east coast, butter clam (*Saxidomus gigantea*) from U.S. west coast, almeja clam (*Venus antiqua*) from Chile, and Atlantic sea scallop (*Plactopecten magellanicus*) from the U.S. east coast. All samples were provided as whole animal homogenates, except Atlantic sea scallop and green mussel, which included hepatopancreas only. Among the naturally contaminated samples, five were blind duplicates tested on separate days that were used for calculation of RSD_r . Samples run as duplicates are indicated in Table 1. Three samples consisting of STX-spiked mussel homogenate (*M. edulis*) at levels that bracketed the regulatory limits of 800 $\mu\text{g}/\text{kg}$ (500 and 1200 $\mu\text{g}/\text{kg}$ spike) and 400 $\mu\text{g}/\text{kg}$ (200 $\mu\text{g}/\text{kg}$ spike) were included to calculate recovery. One sample was the negative control homogenate of *M. edulis* to which the STX spikes were added. All homogenates were extracted by the study participants and the extracts analyzed by RBA in three assays on separate days.

Study Participants

Ten laboratories from seven countries agreed to carry out RBAs for this study, including the United States, Italy, Australia, New Zealand, Thailand, the Philippines, and South Africa. Participants included laboratories from regulatory authorities, as well as government and academic laboratories with monitoring needs. Five of the participating laboratories (Laboratories 1–5) have this method well established and may be considered routine users. Two laboratories had previous experience running this format of the PST RBA, but have not implemented it routinely. One laboratory had previous experience with receptor assays, but had not used the microplate filtration format of the assay. One laboratory had no previous experience with RBAs. Three laboratories from different countries, United States, Chile, and Thailand, carried out the AOAC official mouse bioassay method (AOAC Method 959.08) on the same set of samples. All mouse bioassay laboratories were experienced regulatory authorities with monitoring responsibilities. One laboratory (Health Canada) performed the precolumn oxidation HPLC method for PST (AOAC Method 2005.06).

Preparation of Homogenates

All shellfish samples were thoroughly homogenized using a polytron blender. For spiked samples, saxitoxin standard reference material (STX diHCl) was added to the specified concentration, and the sample was thoroughly rehomogenized to ensure homogeneity. The toxin congener profiles and concentrations of all samples were determined by the precolumn oxidation HPLC method (performed by Health Canada). STX equivalents were determined by mouse bioassay (performed by Maine Department of Marine Resources). Subsamples of each homogenate (12 g) were packaged in polycarbonate tubes and stored at -80°C until shipment to collaborating laboratories by courier. All samples were coded prior to distributing to collaborating laboratories, with the codes to each laboratory being unique, and provided blind. Coding consisted of two letters followed by a number in the form X A1-7, X B1-7, and

X C1-7, where the X indicated the laboratory, the second letter indicated the three assays to be conducted, and the numerical code indicated sample number within that assay. Three practice homogenates were similarly produced.

Shipment of Study Material

The following reagents were provided to the collaborating laboratories in a single shipment containing enough dry ice to keep the contents frozen for 5 days: [³H] STX; STX diHCl standard; rat brain membrane preparation; 21 coded shellfish homogenates; three practice homogenates; and a QC check sample consisting of 18 nM STX diHCl. Sufficient homogenate (12 g) was provided to ensure an accurate weight of material could be removed from the storage vial if an additional extraction were necessary due to unexpected circumstances. The identity of the samples was not released to collaborators. All reagents were received frozen and in good condition. Each participant received electronically a detailed assay protocol, comprehensive instructions for conducting the study and data reporting, and data reporting forms.

Analysis

Participants extracted all homogenates using a modification of the 0.1 M HCl extraction method used in the AOAC standard mouse bioassay protocol (modified only by scale). They were asked to perform three RBAs, each on separate days. Each assay consisted of one 96-well plate that included a standard curve, QC check sample, and seven shellfish extracts. All samples and standards were tested in triplicate wells. All shellfish extracts were run at three dilutions (1/10, 1/50, and 1/200), which ensured that at least one dilution would fall on the linear part of the standard curve. Participants were instructed to analyze samples coded A, B, or C in the first, second, or third assay, respectively, in numerical order. The five blind duplicate samples were coded so that they were tested in two independent assays, with the combination of assays differing between duplicates. Before performing the official study, participants were asked to run a practice assay that included three shellfish homogenates in the same format to ensure that any unexpected problems were encountered and addressed prior to the official study. The practice samples consisted of a negative control mussel homogenate (MLV15), and two naturally contaminated samples that were also included in the full study (MLV05 and MLV11). The identity of the practice samples was not made known to participants. Results of the practice run were submitted by e-mail to the coordinating laboratory for review before proceeding with the full study.

For the mouse bioassay, participants followed the AOAC official mouse bioassay method (AOAC Method **959.08**), with the exception of a modified 0.1 M HCl extraction protocol used in the RBA protocol, which was modified only by scale so that 5 mL 0.1 M HCl was added to 5 g of shellfish homogenate, with all other aspects of the extraction protocol being identical. The HPLC laboratory followed the precolumn oxidation HPLC method for PST (AOAC Method **2005.06**); however, final concentrations in µg/kg and µg STX equiv./kg were calculated using the formula weight of STX diHCl [372 daltons (da)], as opposed to the free base (299.3 da) in the standard HPLC protocol, to more directly compare with the RBA.

Data Analysis and Reporting

Participants were asked to report whether they used a standard or microplate scintillation counter for the study and, if a microplate counter was used, which model, because of differences in inherent counting efficiency between current commercially available counters. For data analysis, participants were instructed to use GraphPad Prism software (La Jolla, CA) or the on-board curve-fitting software provided with their microplate scintillation counter e.g., PerkinElmer Wallac MultiCalc (Gaithersburg, MD) or Packard Top Count software (Packard Instrument Co., Meriden, CT), and to report what software was used. For analysis, a four parameter logistic fit, also known as a sigmoidal dose response with variable slope, or Hill equation, was prescribed. Participants presented their analyzed data on the spreadsheet template provided, including assay quality parameters (slope, IC₅₀, and quantification of the QC check sample), between-well CVs for each sample dilution that fell within the linear part of the standard curve (0.2–0.7 B/B₀), and calculated values for these samples in the well (nM), in the extract (µg STX equiv./mL), and in the shellfish tissue (µg STX equiv./kg). Participants were also asked to report all raw count data so that all results could be analyzed by the coordinating laboratory using identical software (GraphPad Prism 4.0) to assess whether systematic differences in quantification arose from using different curve-fitting software. All data were reported via e-mail to the coordinating laboratory.

The calculated results sheets were reviewed by the coordinating laboratory for obvious errors in dilutions and calculations and for use of the prescribed curve-fitting model. Obvious errors were corrected and the participant laboratory was consulted for concurrence. The reviewed results were then used for evaluation in the collaborative study.

Statistical Evaluation of the Collaborative Study

For each sample analyzed, outliers were first determined using the Grubbs test at a probability value of 1% (www.graphpad.com), with no more than one outlier removed, so that valid data remained from a minimum of eight laboratories. The mean, S_R, and RSD_R, and HorRat values were then calculated for each sample. For blind duplicates, the AOAC INTERNATIONAL Interlaboratory Study Workbook for Blind Duplicates, v2.0, was used to further evaluate for outliers and determine S_r and RSD_r. GraphPad Prism was used to determine correlation among the RBA, mouse bioassay, and HPLC results.

AOAC Official Method 2011.27 Paralytic Shellfish Toxins (PSTs) in Shellfish Receptor Binding Assay First Action 2011

[Applicable to the determination of paralytic shellfish toxins (PSTs), as µg STX diHCl equiv./kg, in shellfish (mussels, clams, scallops) at levels >149 µg STX diHCl equiv./kg, with a limit of detection (LOD) of 45 STX diHCl equiv./kg shellfish and a limit of quantification (LOQ) of 126 µg STX diHCl equiv./kg shellfish.]

Caution: Wear disposable gloves and protective laboratory coat while performing the assay. PSTs are neurotoxins that are harmful if ingested. The assay uses a tritium labeled tracer, [³H] STX, at low concentration.

All laboratories performing the assay must have approved radiation laboratory space and must follow procedures prescribed by their nuclear regulatory agency for receipt, use, and disposal of isotopes.

See Tables 2011.27A–E for results of the interlaboratory study supporting acceptance of the method.

A. Principle

Test portions of shellfish homogenates are extracted using the AOAC mouse bioassay extraction protocol (959.08), modified by scale. The PST receptor assay is a competitive binding assay in which [³H] STX competes with unlabeled STX in standards or mixtures of PST in samples for a finite number of available receptor sites (site 1 on the voltage gated sodium channel) in a rat brain membrane preparation. Following establishment of binding equilibrium at 4°C, unbound [³H] STX is removed by filtration and bound [³H] STX is quantified by liquid scintillation counting. A standard curve is generated using increasing concentrations of STX standard from 10⁻¹⁰ to 10⁻⁶ M STX, which results in a reduction in bound [³H] STX that is directly proportional to the amount of unlabeled toxin present. The concentration of toxin in samples is determined in reference to the standard curve. Incubation is carried out in a microplate format to minimize sample handling and the amount of radioactivity used. Bound [³H] STX (as counts per minute; CPM) can be determined either by conventional or by microplate scintillation counting. Both methods are included in this protocol.

B. Apparatus and Supplies

- (a) *Traditional or microplate scintillation counter.*
 - (b) *Micropipettors.*—1–1000 µL variable volumes and disposable tips.
 - (c) *Eight channel pipettor.*—5–200 µL variable volume and disposable tips.
 - (d) *96-Well microtiter filter plate.*—With 1.0 µm pore size type GF/B glass fiber filter/0.65 µm pore size Durapore support membrane (Millipore, Bedford, MA; Cat. No. MSFB N6B 50).
 - (e) *MultiScreen vacuum manifold.*—Millipore; Cat. No. NSVMHTS00.
 - (f) *Vacuum pump.*
 - (g) *Centrifuge tubes.*—15 and 50 mL, conical, plastic.
 - (h) *Mini dilution tubes in 96-tube array.*
 - (i) *Reagent reservoirs.*
 - (j) *Ice bucket and ice.*
 - (k) *Vortex mixer.*
 - (l) *Sealing tape.*—Millipore; Cat. No. MATA HCL00.
 - (m) *Volumetric flask.*—1 L.
 - (n) *-80°C freezer.*
 - (o) *Refrigerator.*
- For traditional scintillation counter only:
- (p) *MultiScreen punch device.*—Millipore; Cat No. MAMP 096 08.
 - (q) *MultiScreen disposable punch tips.*—Millipore; Cat. No. MADP 196 10.
 - (r) *MultiScreen punch kit B for 4 mL vials.*—Millipore; Cat. No. MAPK 896 0B.
 - (s) *Scintillation vials.*—4 mL.
- For sample extraction:
- (t) *Pipets.*
 - (u) *Centrifuge tubes.*—15 mL, conical, plastic.

- (v) *Vacuum pump or house vacuum.*
- (w) *pH meter or pH paper.*
- (x) *Hot plate.*
- (y) *Graduated centrifuge tubes.*—15 mL.
- (z) *Centrifuge and rotor for 15 mL tubes.*

C. Reagents

- (a) [³H] STX.—0.1 mCi/mL, ≥10 Ci/mmol, ≥90% radiochemical purity (American Radiolabeled Chemicals, St. Louis, MO, or International Isotopes Clearinghouse, Leawood, KS).
 - (b) *STX diHCl.*—NIST RM 8642 (www.nist.gov).
 - (c) *3-Morpholinopropanesulfonic acid (MOPS).*—Sigma (St. Louis, MO; Cat. No. M3183-500G), or equivalent.
 - (d) *Choline chloride.*—Sigma (Cat. No. C7527-500G), or equivalent.
 - (e) *Rat brain membrane preparation.*—See Appendix.
- For traditional counter:
- (f) *Scintiverse BD liquid scintillation cocktail.*—Fisher Scientific (Waltham, MA; Cat. No. SX-18), or equivalent.
- For microplate counter:
- (g) *Optiphase liquid scintillation cocktail.*—PerkinElmer Life Sciences (Downers Grove, IL; Cat. No. 1200-139), or equivalent.
- For sample extraction:
- (h) *Hydrochloric acid (HCl).*—1.0 and 0.1 M.
 - (i) *Sodium hydroxide.*—0.1 M.
 - (j) *Water.*—Distilled or deionized (18 µΩ).

D. Sample Extraction

Accurately weigh 5.0 g tissue homogenate into a tared 15 mL conical tube. Add 5.0 mL of 0.1 M HCl, vortex, and check pH. If necessary, adjust pH to 3.0–4.0 as determined by a pH meter or pH paper. To lower pH, add 1 M HCl dropwise with mixing; to raise pH, add 0.1 M NaOH dropwise with mixing to prevent local alkalization and consequent destruction of toxin. Place the tube in a beaker of boiling water on a hot plate for 5 min with the caps loosened. Remove and cool to room temperature. Check pH and adjust cooled mixture to pH 3.0–4.0 as described above. Transfer entire contents to graduated centrifuge tube and dilute volumetrically to 10 mL. Gently stir contents to homogeneity and allow to settle until portion of supernatant is translucent and can be decanted free of solid particles. Pour approximately 5 to 7 mL of the translucent supernatant into a centrifuge tube. Centrifuge at 3000 × g for 10 min. Retain clarified supernatant and transfer to a clean centrifuge tube. Store extracts at -20°C until tested in receptor assay.

E. Preparation of Stock Solutions and Standards

- (a) *Assay buffer.*—100 mM MOPS/100 mM choline chloride, pH 7.4. Weigh out 20.9 g MOPS and 13.96 g choline chloride and add to 900 mL dH₂O. Adjust pH to 7.4 with NaOH while stirring and bring to a final volume of 1 L with dH₂O. Store at 4°C.
- (b) *Radioligand solution.*—Calculate the concentration of [³H] STX stock provided by the supplier, which may vary between lots. Suppliers generally provide the specific activity in Ci/mmol (generally 10–30 Ci/mmol) and activity in mCi/mL (0.05–0.1 mCi/mL), from which the molar concentration can be calculated. Prepare 4 mL of a 15 nM working stock of [³H] STX fresh daily in 100 mM MOPS/100 mM choline chloride

Table 2011.27A. Receptor binding assay results on individual samples (values are in µg STX diHCl equiv./kg shellfish tissue); summary statistics on all samples; summary statistics excluding Laboratory 9

Assay	No.	Sample															All labs					Labs 1-8		
		Lab															Mean	S _R	RSD _R , %	HorRat	Mean	S _R	RSD _R , %	HorRat
		1	2	3	4	5	6	7	8	9	Mean	S _R	RSD _R , %	HorRat	Mean	S _R	RSD _R , %	HorRat						
Day 1	1	MLV05	370	610	620	410	690	1070	630	660	330	599	222	37.1	2.2	633	212	33.5	2.0					
	2	MLV06	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	422	28.6	1.9	1549	386	24.9	1.7					
	3	MLV08	80	190	140	90	130	160	230	220	100	149	55	37.2	1.8	155	56	36.0	1.7					
	4	MLV09	860	680	950	870	980	1120	1460	820	590	926	255	27.5	1.7	968	237	24.5	1.5					
	5	MLV12	180 ^a	200	200	150	150	100	290	290	100	168	62	37.2	1.8	177	60	34.1	1.7					
	6	MLV14	950	940	1060	1130	1040	750	1460	1320	810	1051	228	21.7	1.4	1081	224	20.7	1.3					
	7	MLV16	660	930	1080	870	840	1320	1490	2420 ^b	490	960	329	34.3	2.1	1027	291	28.3	1.8					
Day 2	8	MLV01	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	818	48.6	3.3	1829	739	40.4	2.8					
	9	MVL02	830	1180	1130	1150	1130	1780	1340	980	690	1134	311	27.4	1.8	1190	281	23.6	1.5					
	10	MLV04	2440	2840	2910	1740	2150	1810	2690	2490	1210	2253	572	25.4	1.8	2384	446	18.7	1.3					
	11	MLV07	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	345	23.9	1.6	1520	279	18.3	1.2					
	12	MLV09	810	1190	1130	810	1630	1390	1880	1120	870	1203	372	30.9	2.0	1245	375	30.1	2.0					
	13	MLV11	270	370	480	340	640	490	240	600	110	393	174	44.3	2.4	429	148	34.4	1.9					
	14	MLV13	400	1240 ^b	560	450	650	530	500	440	200	466	133	28.5	1.6	504	85	16.8	1.0					
Day 3	15	MLV03	330	270	410	180	590	680	370	1570 ^b	90	365	197	54.0	2.9	404	176	43.5	2.4					
	16	MLV05	580	670	250	430	910	700	860	940	300	627	257	41.1	2.4	668	242	36.2	2.1					
	17	MLV06	1290	1520	1460	970	1800	2520	1470	870	1250	1461	488	33.4	2.2	1488	515	34.6	2.3					
	18	MLV07	1010	1600	1390	1000	1720	1860	1520	2150	890	1460	429	29.4	2.0	1531	397	26.0	1.7					
	19	MLV10	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	570	24.0	1.7	2443	569	23.3	1.7					
	20	MLV11	430	350	460	280	550	620	1149 ^b	410	250	419	127	30.2	1.7	443	115	26.0	1.4					
	21	MLV15	ND ^c	ND	ND	ND	ND	ND	ND	180	ND	—	—	—	—	—	—	—	—					
	Avg. RSD _R													33.2					28.7					
	Avg. HorRat													2.0					1.8					

^a CV 41%; not used in calculations.

^b Outlier; not used in calculations.

^c ND = Not detected.

Table 2011.27B. Summary statistics on blind duplicates, run in separate assays (values are in μg STX diHCl equiv./kg)

Lab	MLV05		MLV06		MLV07		MLV09		MLV11		Avg.
	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	
1	370	580	1100	1290	1260	1010	860	810	270	430	
2	610	670	1340	1520	1540	1530	680	1190	370	350	
3	620	250	1320	1460	1220	1390	950	1130	480	401	
4	410	430	1440	970	1980	1000	870	810	340	280	
5	690	910	1260	1790	1760	1720	980	1630	640	550	
6	1070	700	1720	2520	1530	1860	1120	1390	490	620	
7	630	880	2090	1240	1750	1150	1460	1830	230 ^a	1149 ^a	
8	660	940	2130	870	1210	2150	820	1120	600	410	
9	330	300	890	1250	840	890	590	870	110	250	
Avg.		614		1453		1433		1062		416	
S _r		169		432		366		247		83	
S _R		239		444		387		338		152	
RSD _r , %		27.5		29.4		25.5		23.3		20.0	25.1
RSD _R , %		38.9		30.2		27.0		31.9		36.5	32.9
HorRat		2.3		2.0		1.8		2.0		2.0	2.0

^a Outlier; not used in calculation.

buffer. This will provide sufficient volume for one 96-well plate at an in-well concentration of 2.5 nM. Measure total counts of each working stock prior to running an assay: add 35 μL of the working stock [^3H] STX in buffer to a liquid scintillation vial with 4 mL scintillant and count on a traditional liquid scintillation counter. This is done to confirm correct dilution prior to running the assay. Depending on the efficiency of the scintillation counter used, the corresponding CPM will vary, but should be consistent day-to-day and within 15% of the expected value.

(c) *Unlabeled STX standard working solution.*—The STX diHCl standard is provided at a concentration of 268.8 μM (100 $\mu\text{g}/\text{mL}$). A “bulk” standard curve can be made up in advance and stored at 4°C for up to 1 month. The use of a bulk standard curve minimizes the pipetting needed for setting up an assay routinely and improves day-to-day repeatability. Make up 3 mM HCl (e.g., from a 3 M stock, 50 μL in 50 mL), then perform the serial dilutions (see Table 2011.27F) of NIST RM 8642 STX diHCl (100 $\mu\text{g}/\text{mL}$ = 268.8 μM) to make up the standard curve in 3 mM HCl. These standard stock solutions will be diluted 1/6 in the assay to yield the designated in-assay concentrations (see Table 2011.27F).

(d) *Interassay calibration standard (QC check).*—Prepare a reference standard containing 1.8×10^{-8} M STX standard (3.0×10^{-9} M STX in assay) in advance in 3 mM HCl and keep frozen (−80°C) in 1 mL aliquots for long-term storage. Aliquots should be thawed and stored at 4°C for routine use (stable up to 1 month) and analyzed in each assay. This serves as a QC check and confirms day-to-day performance of the assay.

(e) *Rat brain membrane preparation.*—Prepare rat brain membrane preparation in bulk (see Appendix: *Rat Brain Membrane Preparation*) and store at −80°C until used in the assay. Thaw an aliquot of rat brain membrane preparation on ice. Dilute membrane preparation with cold (4°C) 100 mM

MOPS/100 mM choline chloride, pH 7.4, to yield a working stock with a protein concentration of 1.0 mg/mL (this will be diluted in the assay plate to 0.5 mg/mL in-well concentration). Vortex vigorously to achieve a visibly homogeneous suspension. Keep the diluted membrane preparation on ice until ready to use.

F. Performing the Assay

(a) *Plate setup.*—When possible, use a multichannel pipet to minimize pipetting effort and increase consistency. Standard curve, QC check, and sample extracts are run in triplicate wells. Multiple dilutions of sample extracts should be analyzed in order to obtain a value that falls between 0.2–0.7 B/B₀ on the standard curve for quantification. For ease of analysis, it is convenient to use a standard plate layout that maximizes the number of samples and standards that can be analyzed on one plate. For shellfish extracts, a minimum dilution of 1:10 is used, which minimizes potential matrix effects, while still providing an LOQ of approximately 126 $\mu\text{g}/\text{kg}$ shellfish (see Table 2011.27G).

(b) *Addition of samples and standards.*—Add in the following order to each of the 96 wells: 35 μL assay buffer; 35 μL STX standard, QC check, or sample extract; 35 μL [^3H] STX; 105 μL membrane preparation. The assay buffer is added first in order to wet the filter membrane. It is critical to continuously mix the membrane preparation by careful up-and-down pipetting immediately prior to dispensing into the 96-well plate to maintain an even suspension across the entire plate. Cover and incubate plate at 4°C for 1 h.

(c) *Assay filtration.*—Attach the vacuum manifold to the vacuum pump with an in-line side arm flask to catch filtrate from the plate filtration process. Set the vacuum pressure gauge on the pump or vacuum manifold to 4–8” Hg (135–270 millibar), as specified in the instructions provided with the filtration plates. Place the 96-well plate on the vacuum manifold. Fill empty wells with 200 μL MOPS/choline chloride buffer to

Table 2011.27C. Performance of individual laboratories on blind duplicates (values are in µg STX diHCl equiv./kg)

Lab	ID	Day 1	Day 2	Mean	s _r	RSD _r , %	
1	MLV05	370	580	475	148	31.3	
	MLV06	1100	1290	1195	134	11.2	
	MLV07	1260	1010	1135	177	15.6	
	MLV09	860	810	835	35	4.2	
	MLV11	270	430	350	113	32.3	
	Avg.						18.9
2	MLV05	605	670	638	46	7.2	
	MLV06	1340	1520	1430	127	8.9	
	MLV07	1540	1530	1535	7	0.5	
	MLV09	680	1190	935	361	38.6	
	MLV11	370	350	360	14	3.9	
	Avg.						11.8
3	MLV05	620	250	435	262	60.1	
	MLV06	1320	1460	1390	99	7.1	
	MLV07	1220	1303	1262	59	4.7	
	MLV09	950	1130	1040	127	12.2	
	MLV11	480	460	470	14	3.0	
	Avg.						17.4
4	MLV05	410	430	420	14	3.4	
	MLV06	1440	970	1205	332	27.6	
	MLV07	1980	1000	1490	693	46.5	
	MLV09	870	810	840	42	5.1	
	MLV11	340	280	310	42	13.7	
	Avg.						19.2
5	MLV05	690	910	800	156	19.4	
	MLV06	1260	1790	1525	375	24.6	
	MLV07	1760	1720	1740	28	1.6	
	MLV09	980	1630	1305	460	35.2	
	MLV11	640	550	595	64	10.7	
	Avg.						18.3
6	MLV05	1070	700	885	262	29.6	
	MLV06	1720	2520	2120	566	26.7	
	MLV07	1530	1860	1695	233	13.8	
	MLV09	1120	1390	1255	191	15.2	
	MLV11	490	620	555	92	16.6	
	Avg.						20.4
7	MLV05	630	880	755	177	23.4	
	MLV06	2090	1240	1665	601	36.1	
	MLV07	1750	1150	1450	424	29.3	
	MLV09	1460	1830	1645	262	15.9	
	MLV11	230 ^a	1150 ^a				
	Avg.						26.2
8	MLV05	660	940	800	198	24.7	
	MLV06	2130	870	1500	891	59.4	
	MLV07	1210	2150	1680	665	39.6	
	MLV09	820	1120	970	212	21.9	
	MLV11	600	410	505	134	26.6	
	Avg.						34.4

Table 2011.27C. (continued)

Lab	ID	Day 1	Day 2	Mean	s _r	RSD _r , %
9	MLV05	330	300	315	21	6.7
	MLV06	890	1250	1070	255	23.8
	MLV07	840	890	865	35	4.1
	MLV09	590	870	730	198	27.1
	MLV11	110	250	180	99	55.0
Avg.						23.3
Overall avg.						22.2

^a Outlier; not used in calculations.

ensure even vacuum pressure and filtration across the plate. Turn on vacuum. Optimum vacuum will pull the wells to dryness in 2–5 s. Pull contents of all wells through until all liquid is removed. (*Note:* Too low a vacuum will result in slow well clearance, but too high will result in an airlock and no well clearance.) With vacuum pump running, quickly rinse each well twice with 200 µL ice cold MOPS/choline chloride buffer using multichannel pipet. Maintain vacuum until liquid is removed.

(d) *Preparation of the assay for counting.*—Remove the plastic bottom from the plate. Blot the bottom once on absorbent toweling.

(1) *For counting in microplate scintillation counter.*—Place the microplate in a counting cassette. Seal the bottom of the 96-well plate with sealing tape. Add 50 µL Optiphase scintillation cocktail per well using multichannel pipet. Seal the top of the plate with sealing tape. Allow to incubate 30 min at room temperature. Place the plate in a counting cassette and count in a microplate scintillation counter for 1 min/well.

(2) *For counting in traditional scintillation counter.*—Place the microplate in the MultiScreen punch system apparatus. Place the disposable punch tips on top of the microplate. Punch the filters from the wells into scintillation vials and fill with 4 mL scintillation cocktail (Scintiverse or equivalent). Place caps on the vials and vortex. Allow vials to sit overnight in the dark, then count using a tritium window in a traditional scintillation counter.

G. Analysis of Data

For assays performed using the traditional counter, curve fitting is performed using a four-parameter logistic fit, also known as a sigmoidal dose response curve (variable slope; see Figure 2011.27), or Hill equation:

$$y = \min + \frac{\max - \min}{1 + 10^{(x - \log\text{-EC}_{50} \text{ Hill slope})}}$$

where max is the top plateau representing maximum binding in CPM in the absence of competing nonradiolabeled STX, also known as B₀; min is the bottom plateau, equal to nonspecific binding (in CPM) in the presence of saturating nonradiolabeled toxin; IC₅₀ is the inhibitory concentration at which CPM are 50% of max-min (dashed lines; Figure 2011.27); Hill slope is the slope of the curve; x axis is the log concentration of STX; and y axis is total ligand binding in CPM (here represented as B/B₀, or bound/max bound). A curve fitting package such as Prism (GraphPad Software, Inc.) is recommended. For the microplate counter users, receptor

Table 2011.27D. Calibration curve and QC check parameters in three receptor binding assays performed in nine participant laboratories

Lab	Assay day	Slope	IC ₅₀ , nM	QC, nM	Reference, CPM	IC ₇₀ , nM	Standards where RSD >30%; action	Curve fitting software	Scintillation counter	Manual/microplate
1	1	-0.9	1.9	2.4	720	0.90	None	Prism v 3.02	Packard Top Count	Microplate
	2	-1.0	2.0	2.6	733	0.96	None			
	3	-1.1	2.1	3.2	1038	0.92	None			
2	1	-1.1	1.8	3.8	1160	0.66	3 nM; 1 well removed	Prism v 5.0	Packard Top Count	Microplate
	2	-1.2	2.2	3.9	1260	0.85	None			
	3	-1.0	1.6	3.2	1262	0.46	3 nM, 1 nM removed			
3	1	-1.0	2.0	2.3	2529	0.41	First column removed	Prism v 5.0	Wallac Microbeta	Microplate
	2	-0.9	2.0	2.5	1463	0.92	1000 nM; 1 well removed			
	3	1.0	1.6	2.8	2088	0.80	None			
4	1	-0.9	1.7	3.4	1125	0.61	None	Prism v 3.03	PerkinElmer Tricarb	Manual
	2	-1.2	1.7	3.2 ^a	1611	0.77	None			
	3	-0.9	1.2	2.9	1324	0.45	30 nM 35%; 1 well removed			
5	1	-0.9	1.4	3.3	1566	0.64	1.0 nM; 1 well removed	MultiCalc	Wallac Microbeta	Microplate
	2	-1.2	1.8	3.6	1528	1.05	0.1 nM and 30 nM; 1 well removed			
	3	-1.2	1.8	2.9	1052	0.67	None			
6	1	-1.1	2.6	3.0	670	1.15	None	Prism v 4.0	Wallac Microbeta	Microplate
	2	-1.0	2.0	4.0 ^b	1124	1.08	None			
	3	-1.1	3.4	6.5 ^b	1030	2.04 ^c	None			
7	1	-0.8	1.0	2.8 ^a	919	0.33	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	1.6	2.7	619	0.70	None			
	3	-0.9	2.1	3.2 ^a	693	0.82	None			
8	1	-1.2	1.7	3.7	1146	0.86	None	Prism	Wallac Microbeta	Microplate
	2	-1.1	1.4	1.5 ^b	1095	0.78	None			
	3	-1.1	2.4	2.3	886	1.04	None			
9	1	-1.0	2.2	4.0 ^b	1363	0.97	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	2.0	3.2	1380	0.85	100 nM 33%; left in			
	3	-1.0	2.1	3.7	1532	0.92	None			

^a One well removed.

^b Outside of specifications.

^c Outlier by Grubbs test.

assay applications provided by the manufacturer may be used (e.g., MultiCalc; PerkinElmer Wallac, Gaithersburg, MD).

(a) *Sample quantification.*—Sample quantification is carried out only on dilutions that fall within B/B₀ of 0.2–0.7, where B represents the bound [³H]STX (in CPM) in the sample and B₀ represents the max bound [³H]STX (in CPM). Where more than one dilution falls within B/B₀ of 0.2–0.7 on the curve, all sample wells corresponding to these dilutions are used to calculate sample concentration. Sample concentration is calculated in µg STX diHCl equiv./kg shellfish, from the in-well nM concentration obtained from the curve fitting software using the following formulas:

$$(nM \text{ STX equiv}) \times (\text{sample dilution}) \times \frac{(210 \mu\text{L total volume})}{35 \mu\text{L sample}} = nM \text{ STX equiv in extract}$$

$$(nM \text{ STX diHCl equiv. in extract}) \times \frac{1 \text{ L}}{1000 \text{ mL}} \times \frac{372 \text{ ng}}{\text{nmol}} \times \frac{1 \mu\text{g}}{1000 \text{ ng}} = \mu\text{g STX diHCl equiv./mL}$$

$$\mu\text{g STX diHCl equiv./mL} \times \frac{\text{mL extract}}{\text{g shellfish}} \times \frac{1000 \text{ g}}{\text{kg}} = \mu\text{g STX diHCl equiv./kg}$$

H. Assay Performance Standards

The following criteria must be met for assay acceptance:

Table 2011.27E. Results of the receptor binding assay (RBA), mouse bioassay (MBA), and HPLC analyses of 21 shellfish extracts, sorted by mouse bioassay value (all values are in µg STX diHCl equiv./kg shellfish tissue; results in bold indicate toxicity above the 800 µg STX diHCl equiv./kg regulatory limit; all other results indicate toxicity below the regulatory limit)

Sample	Lab 1	Lab 2	Lab 3	Lab 4	Lab 5	Lab 6	Lab 7	Lab 8	Lab 9	RBA, avg.	HPLC	MBA
21	ND ^a	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
5	180	200	200	150	150	100	150	290	100	168	108	ND
15	330	270	410	180	590	680	370	1570^b	90	365	196	182
13	270	370	480	340	640	290	240	600	110	371	236	299
20	430	350	460	280	550	490	1150^b	410	250	403	236	299
14	400	1240^b	560	450	650	530	500	440	200	466	625	343
1	370	610	620	410	690	1070^b	630^b	660	330	599	413	387
16	580	670	250	430	910	700	860^b	940^b	300	627	413	387
3	80	190	140	90	130	160	230	220	100	149	341	405
6	950	940	1060	1130	1040	750	1460	1320	810	1051	618	485
7	660	930	1080	870	840	1320	1490	2420	490	960	685	528
2	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	931	595
17	1290	1520	1460	970	1800	2520	1470	870	1250	1460	931	595
4	860	680	950	870	980	1120	1460	820	590	926	1070	653
12	810	1190	1130	810	1630	1390	1880	1120	870	1203	1070	653
11	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	965	714
18	1010	1600	1390	1000	1720	1860	1520	2150	890	1452	965	714
8	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	894	752
9	830	1180	1130	1150	1130	1780	1340	980	690	1134	802	792
19	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	2000	1027
10	2440	2840	2910	1740	2150	1800	2690	2490	1210	2252	1890	1080

^a ND = Not detected.

^b Outlier; not used in average calculation.

(a) For a ligand that specifically binds at one receptor site, the slope of the resulting competition curve should theoretically be -1.0. If the slope of the curve for a given assay is outside of the acceptable range of -0.8 to -1.2, linearity of the assay will be compromised and quantification of the unknowns will be incorrect.

(b) RSDs of triplicate CPMs for standards should be below 30% as variability may affect the slope calculation and thereby quantification of samples.

(c) If the IC₅₀ is out of the acceptable range (2.0 nM ± 30%) then the assay should be considered suspect and rerun, as a shift in the curve will result in over- or underestimation of sample concentrations.

(d) QC check should be 3 nM STX ± 30% (in-well concentration). Assays with a QC check sample out of specifications should trigger a check of the IC₅₀ value.

The following criteria must be met for acceptability of a sample measurement:

(a) Sample quantification should be done only on dilutions that fall within B/B₀ of 0.2–0.7. In the event that all sample dilutions fall below B/B₀ 0.2 (i.e., concentration is too high), further dilutions must be made and the sample reanalyzed. In the event that the sample concentration is too low to be quantified (i.e., B/B₀ > 0.7), the sample is reported as below LOD. If more

Table 2011.27F. Dilution series to prepare bulk solutions for standard curve

	Stock, M	In-assay, M
100 µL 268.8 µM STX + 4.38 mL 0.003 M HCl	6 × 10 ⁻⁶	1 × 10 ⁻⁶
500 µL 6 × 10 ⁻⁶ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁷	1 × 10 ⁻⁷
1.5 mL 6 × 10 ⁻⁷ M + 3.5 mL 0.003 M HCl	1.8 × 10 ⁻⁷	3 × 10 ⁻⁸
500 µL 6 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁸	1 × 10 ⁻⁸
500 µL 1.8 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	1.8 × 10 ⁻⁸	3 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁸ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁹	1 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁹ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻¹⁰	1 × 10 ⁻¹⁰
5 mL 0.003 M HCl	0	Reference

Table 2011.27G. Recommended microplate layout for ease of handling triplicate wells of standard curve, QC check sample, and unknown samples; each sample is run at three dilutions (1:10, 1:50, 1:200); standard curve is run in columns 1–3 (values are in M STX)^a

Microplate row	Microplate column											
	1	2	3	4	5	6	7	8	9	10	11	12
A	10 ⁻⁶	10 ⁻⁶	10 ⁻⁶	QC	QC	QC	U3	U3	U3	U6	U6	U6
							1:50	1:50	1:50	1:10	1:10	1:10
B	10 ⁻⁷	10 ⁻⁷	10 ⁻⁷	U1	U1	U1	U3	U3	U3	U6	U6	U6
				1:10	1:10	1:10	1:200	1:200	1:200	1:50	1:50	1:50
C	3 × 10 ⁻⁸	3 × 10 ⁻⁸	3 × 10 ⁻⁸	U1	U1	U1	U4	U4	U4	U6	U6	U6
				1:50	1:50	1:50	1:10	1:10	1:10	1:200	1:200	1:200
D	10 ⁻⁸	10 ⁻⁸	10 ⁻⁸	U1	U1	U1	U4	U4	U4	U7	U7	U7
				1:200	1:200	1:200	1:50	1:50	1:50	1:10	1:10	1:10
E	3 × 10 ⁻⁹	3 × 10 ⁻⁹	3 × 10 ⁻⁹	U2	U2	U2	U4	U4	U	U7	U7	U7
				1:10	1:10	1:10	1:200	1:200	1:200	1:50	1:50	1:50
F	10 ⁻⁹	10 ⁻⁹	10 ⁻⁹	U2	U2	U2	U5	U5	U5	U7	U7	U7
				1:50	1:50	1:50	1:10	1:10	1:10	1:200	1:200	1:200
G	10 ⁻¹⁰	10 ⁻¹⁰	10 ⁻¹⁰	U2	U2	U2	U5	U5	U5			
				1:200	1:200	1:200	1:50	1:50	1:50			
H	REF	REF	REF	U3	U3	U3	U5	U5	U5			
				1:10	1:10	1:10	1:200	1:200	1:200			

^a REF = Reference; QC = quality control check; U = unknown sample. [Note: The same standard curve may be used for multiple plates (i.e., 11 samples can be run on subsequent plates in a series if the standard curve is not included).]

than one dilution falls on the linear part of the curve, an average value calculated from all dilutions should be used. If there is disagreement between different dilutions in final concentration reported, check for error in the sample dilution process.

(b) RSD of the sample CPMs should be ≤30%.

Reference: *J. AOAC Int.* **95**, 795(2012)

Results and Discussion

Sample Characterization

All shellfish homogenates (MLV1–16) were analyzed by

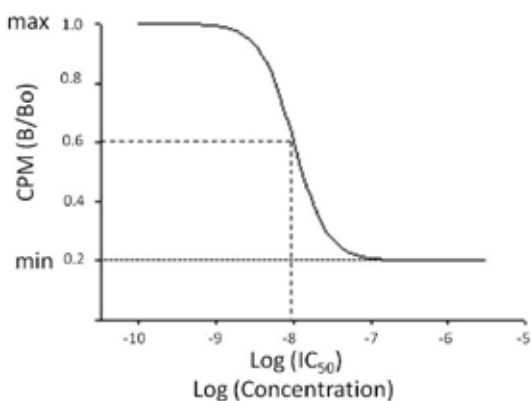


Figure 2011.27. Sigmoidal dose response curve. Dashed lines indicate log IC₅₀.

HPLC using the precolumn oxidation method (AOAC Method **2005.06**) to determine toxin congener profiles and quantify total PST as µg STX diHCl equiv./kg prior to initiation of the study (Table 2). It is noteworthy that the clear majority of samples, irrespective of shellfish species and location, were dominated largely by STX and GTX2,3 whereas the N1-hydroxylated congeners NEO and GTX1,4 were virtually absent, except in blue mussel from the U.S. west coast. The most unusual profile was observed in green mussel, which was dominated by the weakly toxic N-sulfo-carbamoyl congeners C1,2. The samples were analyzed by the AOAC mouse bioassay (AOAC Method **959.08**) by three laboratories that routinely perform the mouse bioassay for regulatory purposes (Table 3). The mouse bioassay detection limit is approximately 400 µg STX diHCl equiv./kg (one laboratory reported values as low as 290 µg STX equiv./kg). Because the study design included samples that bracketed the lower regulatory limit of 400 µg STX diHCl equiv./kg, several samples were reported as being below the mouse bioassay detection limit. For samples in which all values were above the detection threshold, the between-laboratory RSD_R of the mouse bioassay was 18.9%.

Data Reporting and Initial RBA Data Review

Nine of the 10 laboratories that received the study materials completed the study and reported results. All nine carried out the practice assay and reported results to the coordinating laboratory, which evaluated the results and provided feedback to the participating laboratories before initiating the full study. Following completion of the full study, the participating laboratories provided all raw and calculated data for each of

the three assays performed via e-mail to the coordinating laboratory. The calculated results sheets were reviewed by the coordinating laboratory for obvious errors in sample dilutions and calculations, and for the use of the prescribed curve-fitting model. One laboratory used a sigmoidal curve-fitting model with the slope set to 1 (one-site binding curve in Prism), rather than the prescribed four-parameter logistic fit. In this case, the raw data were reanalyzed by the coordinating laboratory using the prescribed method. Obvious errors in calculation were corrected, such as accounting for the two-fold sample dilution resulting from the extraction process. In some cases, the participating laboratory carried out a fourth assay due to variability or inconsistency among dilutions for selected samples. In these cases, the value reported from the repeat (fourth) assay was used. One laboratory had consistent disagreement between the 1/50 and 1/200 dilutions when both fell within B/B_0 , 0.2–0.7. In all cases the 1/200 dilution overestimated almost two-fold relative to the 1/50 dilution, suggesting a systematic dilution error. In standard practice, these samples should be rerun. However, the instructions did not direct the participants to do so. Therefore, where there was corroborative evidence for the value reported by the 1/50 dilution, based on the 1/10 dilution, the 1/200 dilution was omitted. Where there was no basis on which to exclude the 1/200 value, an average value was calculated. This tended to result in an overestimate, and in two cases resulted in statistical outliers.

Overall Performance of the Method: Reproducibility

Table 2011.27A summarizes the results obtained for 21 individual shellfish samples analyzed in three RBAs, determined by nine participating laboratories. Samples 1–7 were analyzed in the first assay, samples 8–14 in the second assay, and samples 15–21 in the third assay. Among these samples were five blind duplicates, treated here as individual unknown samples. One sample (marked by an footnote a in Table 2011.27A) had a high variability in CPM between wells that was not attributable to any known cause, and was, therefore, omitted from analysis. Outliers identified by Grubbs test ($P < 0.01$) were excluded from the analysis (marked by footnote b in Table 2011.27A). The overall RSD_R among all 21 independent samples was 33.2%, resulting in an average HorRat value of 2.0 (Table 2011.27A). The HorRat values on individual samples ranged from 1.4 to 3.3, with a median value of 1.8. There was no apparent trend in reproducibility according to sample concentration or among shellfish species. If only the laboratories that are routine users of the RBA for PST (Laboratories 1–5) are included in the analysis, the average RSD_R is 23.1%, resulting in an average HorRat value of 1.4. Laboratory 9 tended to report the lowest values among the participating laboratories (14 of 21 samples), and although its individual sample values were not found to be statistical outliers, removing the results of this laboratory reduces all but one HorRat value (which remains unchanged), yielding an average HorRat value of 1.8 (range 1.0–2.8; Table 2011.27A). Removal of any other single laboratory's results does not appreciably change the overall study performance. The reason for the systematically low values reported by Laboratory 9 is not clear, since the assay parameters fall well within those reported by the other laboratories. Given that assay parameters are within normal range, one possible source of systematic error

could be incomplete extraction or pH adjustment of extracts, either of which would result in lower toxicity values.

A comparison of the RBA reproducibility with that of existing AOAC *Official Methods* is instructive. The AOAC collaborative study of the mouse bioassay (11), which entailed the analysis of seven samples representing three levels of STX-spiked shellfish by 11 participating laboratories, yielded a similar average RSD_R of 22%. More recent proficiency tests of the mouse bioassay performed in European regulatory laboratories report RSD_R of 2.3–38.3% on three samples run by eight laboratories (2) and RSD_R of 18.1–44.8% on two samples run by 20 laboratories (12). The mouse bioassay RSD_R values obtained in the current study ranged from 1.1 to 46.3% (average 19%) for three laboratories. The collaborative studies of the HPLC methods report reproducibility values for individual PST congeners, but do not report reproducibility of the composite toxic potency values. Collaborative study of the precolumn oxidation HPLC method (AOAC Method 2005.06) resulted in an average RSD_R of 27.0% and HorRat value of 1.3 (range 0.8–2.1) for STX following C18 cleanup, but the reproducibility of other congeners varied considerably, with the maximum HorRat value (4.7), exceeding the highest HorRat value obtained by RBA (3.3).

Because composite toxic potency values were not reported in the studies of the HPLC methods, it is uncertain how this variability influences the composite toxic potency calculated from these methods. The average and ranges of HorRat values obtained for different congeners were: neoSTX–1.7 (range 1.2–2.5); dcSTX–1.1 (range 0.6–2.1); GTX1,4–1.9 (range 1.1–4.2), GTX2,3–1.4 (range 0.8–1.9); B1–1.1 (range 0.7–1.9); and C1,2–1.6 (range 0.9–4.5). Because of the variability obtained in neoSTX, GTX1,4, C3,4, and B2, AOAC Method 2005.06 calls for a second SPE-COOH cleanup of samples suspected of containing these congeners, after which reproducibility improved somewhat: neoSTX–1.8 (range 1.3–2.1); GTX1,4–1.3 (range 1.0–2.1); and C3,4–1.2 (range 0.8–1.8). The postcolumn oxidation HPLC method (AOAC Method 2011.02) reported an average HorRat value of 0.6 for STX. In this method, neoSTX with an average HorRat of 1.9 (range 0.6–4.0) and GTX4 with an average HorRat of 1.6 (range 1.0–2.9) had reproducibility values that may affect the overall composite potency values. The maximum HorRat value (4.0) reported in this study also exceeded the maximum value reported in the RBA.

In summary, with the removal of Laboratory 9, the overall reproducibility of the RBA falls within the performance measures achieved by the established AOAC *Official Methods* for PST. The difference in reproducibility achieved by the laboratories that are routine users of the assay and participants who are not routine users of the method highlights the importance of training if this method were to be implemented in a regulatory setting.

Within-Laboratory Repeatability

Within-laboratory variability (RSD_r) was determined on five samples that were provided as blind duplicates. Participants were unaware that blind duplicates were included among the coded samples received. The duplicate samples were coded so that they were analyzed in separate assays, with different duplicate pairs falling into different assays (Table 1). One outlier was found among the results of the blind duplicates by Cochran's

test, $P < 0.025$ (Laboratory 7, sample MLV11) using the AOAC INTERNATIONAL Interlaboratory Study Workbook for Blind Duplicates, v2.0. An overall RSD_r of 25.1% was observed, with an RSD_R of 32.9%, yielding a HorRat value of 2.0, similar to that of the overall study (Table 2011.27B). When the performance of individual laboratories was evaluated separately, the average RSD_r was 22.2%, with individual laboratories varying from 11.8 to 34.4% (Table 2011.27C). Routine users of the microplate format of the PST RBA (Laboratories 1–5) obtained an average RSD_r of 17.1%, which is similar to that obtained in the SLV study (10), and lower than that obtained by nonroutine users (Laboratories 6–9), which averaged 26.1% and ranged as high as 34.4%. The AOAC collaborative study of the mouse bioassay (11) did not report RSD_r ; however, analysis of the data from that study using AOAC INTERNATIONAL's Interlaboratory Study Workbook for Blind Duplicates results in an average RSD_r of 16.5% for three STX-spiked samples. Proficiency testing of the mouse bioassay performed in eight French laboratories reported an average RSD_r of 8.3% on three samples (2). The analysis of blind duplicates in the collaborative study of the precolumn oxidation HPLC method (AOAC Method 2005.06) resulted in an RSD_r of 15.2% for STX following SPE C18 cleanup and an average RSD_r of 16.4% across all congeners, which ranged from 6.0 to 31.7%. Following SPE-COOH cleanup, repeatability was similar, with RSD_r of 17.2% across all congeners. The intralaboratory repeatability values obtained in the postcolumn oxidation HPLC method (AOAC Method 2011.02) averaged 6.4% for STX; most other congeners were similar, with neoSTX being the only congener that showed a somewhat higher RSD_r of 23.3%.

In summary, the within-laboratory repeatability of the RBA was found to be acceptable, with all but two laboratories achieving an RSD_r of 23.3% or less, and the routine users of the assay achieving an average RSD_r of 17.1%.

Spike Recovery

Three samples included in the study were homogenates of blue mussel spiked with STX diHCl at concentrations intended to bracket the regulatory limits of 800 μg STX equiv./kg used by most countries and 400 μg STX equiv./kg imposed in the Philippines. Nominal concentrations in the spiked samples were 200, 500, and 1200 μg STX equiv./kg. Also included in the study was the blue mussel homogenate to which the STX spikes had been added, which was determined to be negative for STX by the precolumn oxidation HPLC method. The negative control homogenate was reported as nondetectable by eight of nine laboratories. Recovery of spiked STX by the RBA was 84.4, 93.3, and 88.1%, respectively, for the 200, 500, and 1200 μg STX diHCl equiv./kg spike levels, and yielded a slope of 0.87 and r^2 of 0.86 (Figure 2). In the current study, the mouse bioassay reported < detection limit, and 68.6 and 40.5% recovery for the 200, 500, and 1200 μg STX diHCl equiv./kg spike levels. The AOAC collaborative study of the mouse bioassay (11) reported recoveries of 62.3% at spike levels similar to those in the current study (equivalent to 1000 μg STX diHCl equiv./kg) but higher recoveries of 81.5 and 96.0% were achieved at higher spike levels equivalent to 4000 and 8000 μg STX diHCl equiv./kg.

The observed poor recovery in the mouse bioassay at concentrations near and below the regulatory limit has been observed in other studies (2), and has been attributed to a

salt or protective effect of the shellfish matrix, which, for concentrations at or below the regulatory limit of 800 $\mu\text{g}/\text{kg}$, is injected undiluted into the mouse. The spike recovery observed in the precolumn HPLC method in this study is also somewhat low, with 54.0, 62, and 51.5% recovery at the 200, 500, and 1200 μg STX diHCl equiv./kg spike levels, respectively. The AOAC collaborative study of the precolumn HPLC method reported 74.4–76.8% at similar spike levels following SPE C18 cleanup and 63.7–68.2% following SPE-COOH cleanup (3, 4). In comparison, the postcolumn HPLC method reported 88–104% recovery of STX spiked at levels somewhat lower than the current study. The higher recovery of the RBA than the HPLC method in the current study may reflect the use of the 0.1 M HCl extraction method in the RBA as compared to the acetic acid extraction used in the HPLC methods.

We previously established in the SLV study that the RBA performs well with shellfish extracted using either method (10). In that study, the RBA reported slightly higher toxicity values for shellfish extracts made using the 0.1 M HCl method than the acetic acid extraction, yielding a correlation of 0.99 with a slope of 1.23 (10). The higher toxicity reported by the RBA in 0.1 M HCl extracts may reflect the hydrolysis of less toxic congeners to more toxic congeners.

Assay Parameters and Quality Metrics

Table 2011.27D summarizes the assay parameters and quality metrics for all laboratories. Eight of nine laboratories used microplate scintillation counters. Laboratory 4 used the manual counting method in which the microplate well filters are punched out, using an eight-place punch system, into traditional 4 mL scintillation vials and counted. Its performance using the manual counting method (RSD_r 17.4%) was similar to or better than that of the laboratories using the microplate method, indicating that using the manual counting method does not affect the performance of the assay. Similarly, there was no apparent difference in assay parameters when the Packard Top Count (single detector) was used, compared to the Wallac Microbeta (coincidence detector), although the reference CPM values obtained on the Top Count generally were somewhat lower due to differences in counting efficiency inherent in the differences in detector geometry. Eight of nine laboratories used GraphPad Prism for curve-fitting, while only Laboratory 5 used Wallac MultiCalc software. Values reported by Laboratory 5 fell well within the range of values reported by laboratories using Prism.

All assays resulted in slopes between -0.8 and -1.2 , as specified in the protocol. This specification reflects the fact that in a competitive binding assay for a ligand that interacts specifically at a single receptor site, the slope of the resulting standard curve should theoretically be 1.0. Although curve-fitting software packages often include a one-site binding curve that fixes the slope at 1.0, we specified in the protocol the use of the four-parameter logistic fit (also known as sigmoidal dose-response with variable slope), because it more readily identifies problems with the standard curve that may skew results. Laboratory 9 reported results using a one-site binding curve fit; in this case, the coordinating laboratory recalculated their raw data using the four-parameter logistic fit. The protocol also calls for $RSD\% < 30$ on all standards. Most analysts did not experience variability problems in the standard wells. Infrequent high RSD s were most often associated with the well

Table 2. Congener profiles in shellfish homogenates included in the collaborative study^a

Sample name	Species	STX	NEO	dcSTX	GTX1,4	GTX2,3	dcGTX2,3	B1	C1,2	C3,4	Total PSP	µg STX diHCl equiv./kg
MLV01	Surf clam	639.8		74.0		226.2	207.0				1146.9	894.3
MLV02	Almeja clam	298.3				1290.1		266.6			1855.0	802.1
MLV03	Chorito mussel	77.6				310.4					388.0	195.5
MLV04	Atlantic sea scallop	831.6				2785.6					3617.3	1890.2
MLV05	Atlantic sea scallop	193.8				576.2					770.0	412.8
MLV06	California mussel	912.8		10.9		0.0		233.8			1157.5	931.3
MLV07	Blue mussel, U.S. east coast	548.2				1097.3					1645.5	965.2
MLV08	Green mussel	164.2		63.5			272.3	454.8	3629.0		4419.6	340.8
MLV09	Blue mussel, U.S. west coast	432.3	124.9	8.7	353.7	727.8		506.4			2153.9	1070.9
MLV10	Butter clam	1763.5		40.6		533.2		203.5			2540.8	2000.9
MLV11	Almeja clam	159.1		12.2		185.5					356.8	236.9
MLV12	Blue mussel spike	108.4									108.4	108.4
MLV13	Blue mussel spike	310.2									310.2	310.2
MLV14	Blue mussel spike	618.5									618.5	618.5
MLV15	Blue mussel blank										0.0	0.0
MLV16	Chorito mussel	389.8		14.3		754.1					1158.1	684.9

^a Values for individual congeners are in µg/kg. Values for composite toxicity are in µg STX diHCl equiv./kg. Abbreviations for congeners are as follows: STX – saxitoxin; NEO – neosaxitoxin; dcSTX – decarbamoyl saxitoxin; GTX1,4 – gonyautoxin 1 and gonyautoxin 4; GTX2,3 – gonyautoxin 2 and gonyautoxin 3; B1 – gonyautoxin 5 (also known as sulfocarbamoyl STX B1); C1,2 – sulfocarbamoyl STX C1 and sulfocarbamoyl STX C2; C3,4 – sulfocarbamoyl STX C3 and sulfocarbamoyl STX C4.

Table 3. Mouse bioassay results on collaborative study samples from three laboratories^a

Sample No.	Sample ID	MBA Lab A	MBA Lab B	MBA Lab C	MBA Avg.	MBA s _R	MBA RSD _R , %
1	MLV05	400	415	340	385	39.7	10.3
2	MLV06	550	597	540	562	30.4	5.4
3	MLV08	440	<dl ^b	370	405	49.5	12.2
4	MLV09	670	612	760	681	74.6	11.0
5	MLV12	<dl	<dl	<dl	—	—	—
6	MLV14	489	489	480	486	5.2	1.1
7	MLV16	585	585	470	547	66.4	12.1
8	MLV01	750	716	600	689	78.6	11.4
9	MLV02	670	1115	590	792	282.9	35.7
10	MLV04	2040	<dl	1080	1560	678.8	43.5
11	MLV07	1480	748	670	966	446.8	46.3
12	MLV09	—	594	670	602	11.3	1.9
13	MLV11	380	379	<dl	380	—	—
14	MLV13	<dl	343	<dl	343	—	—
15	MLV03	400	364	<dl	382	—	—
16	MLV05	—	396	370	383	18.4	4.8
17	MLV06	—	702	630	666	50.9	7.6
18	MLV07	—	<dl	690	690	—	—
19	MLV10	1320	890	870	1027	254.2	24.8
20	MLV11	—	364	290	327	52.3	16.0
21	MLV15	<dl	<dl	<dl	—	—	—

^a Values are in µg STX diHCl equiv./kg.

^b dl = Detection limit.

in column 1 of the 96-well plate. Most analysts removed the suspect well from the curve-fitting process. When the RSD for a given standard was near the stated cutoff (e.g., 31–33%), and left in the curve-fitting process, there was no apparent effect on the curve parameters listed as criteria for assay acceptance.

The average IC₅₀ among all 27 assays was 1.9 + 0.45 nM (RSD_R 23.5%). The other assay quality metric called for by the protocol is the analysis of the QC check sample, which should be 3 ± 0.9 nM STX (30% RSD, in-well concentration). Four of the 27 assays had QC values outside the stated limits, with no obvious error responsible for the variability. Among these, Laboratory 7 reported 6.5 nM for the QC check in assay 3 and an IC₅₀ of 3.4 nM, which was outside the norm. Similarly, Laboratory 8 reported a QC of 1.5 nM in assay 2 and a low IC₅₀ of 1.4 nM, which is at the lower edge of acceptability. In general practice, these values would trigger repeating the assay. However, because of the minimal number of laboratories participating in the study, both of these assays were retained in the study. In neither case were the reported sample values systematically higher or lower than those reported in the other assays.

LOD and LOQ

The LOD was calculated based on the measurement of the negative control shellfish matrix (MLV15) using the blank + 3×SD approach according to Eurachem guidelines (13), as

recently applied to AOAC Method **2006.02**, an ELISA for domoic acid in shellfish using a similar four-parameter logistic curve (14). All laboratories reported <dl for this sample using the prescribed cutoff of B/B₀ <0.7 for quantification, with the exception of Laboratory 8, which was removed as an outlier as determined by Grubbs test ($P < 0.01$). If these samples are instead quantified using the B/B₀ values obtained, a mean of 5.5 ng/mL is obtained with an SD of 5.7 ng/mL, resulting in an LOD of 45 µg STX diHCl equiv./kg. Using the blank + 10×SD definition, an LOQ of 126 µg STX di HCl equiv./kg is thus obtained. We previously established empirically that a 1/10 dilution of shellfish extracts is sufficient to remove matrix effects in the RBA (10), when a quantification cutoff of B/B₀ <0.7 is used. This is the basis for the ten-fold minimum sample dilution used in the current study. The IC₇₀ values (B/B₀ 0.7) for all standard curves run in the study are presented in Table **2011.27D**. An average of 0.80 ± 0.188 nM STX diHCl was obtained across all assays, following the removal of one outlier based on the Grubbs test ($P < 0.01$). Applying the blank + 3×SD to this value, an LOD of 64 µg STX diHCl equiv./kg is obtained; applying the blank + 10×SD to this value results in an LOQ of 131 µg STX diHCl equiv./kg for a sample diluted 1/10 and extracted as indicated in the study, in fair agreement with the value calculated above.

Correlation with HPLC and Mouse Bioassay

Comparison of the RBA results with the mouse bioassay

Nominal	Avg	S _R	RSD _R , %	Recovery, %
200	169	58	34.6	84.4
500	466	133	28.5	93.3
1200	1057	228	21.7	88.1

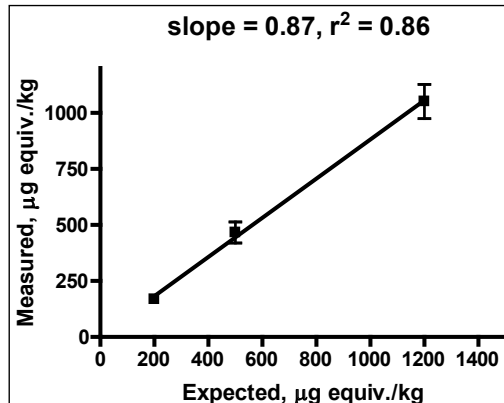


Figure 2. Recovery of spiked STX diHCl in homogenates of blue mussel. Values are in µg STX diHCl equiv./kg.

results yielded an r^2 of 0.84 and a slope of 1.64, indicating that the RBA reports somewhat higher STX equivalents in shellfish, relative to the mouse bioassay (Figure 3). This overestimate has been previously reported for both RBA and HPLC methods (2, 9) at the STX levels near or below the regulatory limit, which are the focus of the current study. Consistent with these findings, the HPLC method also reported higher values than the mouse bioassay in this study, with a slope of 1.33 and an r^2 of 0.84. RBA results correlated better with the precolumn oxidation HPLC method, with a slope of 1.20 and an r^2 of 0.92.

RBA Yielded No False Negatives Relative to the Regulatory Limit

When the data from the three methods were sorted by increasing µg STX diHCl equiv./kg as reported by the mouse bioassay, the RBA did not report any false negatives when compared to the regulatory limit of 800 µg STX equiv./kg (Table 2011.27E). When compared with the precolumn oxidation HPLC method, only Laboratory 9 reported values lower than the HPLC method. The fact that the RBA reports somewhat higher toxicity than the mouse bioassay or HPLC at levels near or below the regulatory limit is beneficial from a food safety standpoint. The higher values reported presumably arise from better recoveries, as demonstrated above. From a shellfish producer's perspective, the improved detection limits relative to the mouse bioassay and better recovery of low toxin levels compared to the HPLC can help to provide advance warning of developing toxicity, allowing producers to harvest early, delay harvest, or move cultures, as appropriate.

Participants' Comments

Laboratory 6 participated in the study without previous

experience running receptor assays, and in doing so, identified several points needing clarification that have since been added to the proposed *Official Method* as enumerated in this report: (1) The vacuum required for filtration was not specified at 4–8" Hg, which is critical because insufficient vacuum pressure results in too slow a clearance of the wells, whereas too much pressure results in an airlock and no filtration at all. (2) Scintillation counting time for the microplates is 1 min/well. (3) Instructions have been added regarding how to calculate sample concentration if more than one dilution falls within B/B₀ 0.2–0.7; specifically, an average value should be calculated from all sample dilutions falling within B/B₀ 0.2–0.7. When corrected for dilution, serial sample dilutions should yield similar quantification. The absence of linearity between sample dilutions indicates either error in dilution or sample matrix interference; however, at the minimum sample dilutions recommended in the proposed *Official Method*, matrix effects from shellfish homogenates have not been encountered (10). In the current study, the nonlinearity of dilutions experienced in several samples by Laboratory 8 was not observed by the other laboratories, suggesting a systematic sample dilution issue rather than a sample matrix problem. Although experienced in RBAs in general, Laboratory 8 had not previously run the microplate filtration format of the assay for PST.

Laboratory 9, which reported generally lower values than the other laboratories, although familiar with the assay, had not performed it in more than a year. The lower values reported do not appear to be associated with conduct of the assay, or scintillation conduct of the assay, or scintillation counting, since the assay metrics are well within the averages reported by the other laboratories. Insufficient boiling or pH adjustment of sample extracts are a possible explanation. These points identified by the study participants should be added to the critical steps identified in the SLV study (10) that can affect precision and accuracy of the assay results, including: (1) ensure that the water is strongly boiling during extraction; (2) carefully adjust pH of extracts; (3) ensure even distribution of the membrane preparation across the microplate by frequent vortex-mixing or pipetting before and during its addition to the plate; (4) the wells must clear within 2–5 s during filtration; (5) the wash buffer should be ice-cold to minimize the rate of toxin release from the receptor; and (6) following addition of scintillant to the wells, incubate a minimum of 30 min to ensure that the scintillant fully penetrates the filters before counting.

Recommendations

The collaborative study of the RBA for PST was completed by nine laboratories representing six countries. Collaborators quantified PST as a composite toxicity value reported in µg STX di HCl equiv./kg in a variety of shellfish species from different regions of the world, containing varied toxin congener profiles. The study included laboratories with extensive experience as well as others with little or no previous experience. The study also included both microplate and scintillation counters as end points, because either instrument type could potentially be used by test laboratories. The study demonstrates that the RBA yields adequate repeatability, reproducibility, and recovery for routine determination and monitoring of PST in shellfish. The greater precision attained by laboratories that received prior training on the RBA and routinely implement this assay suggests that

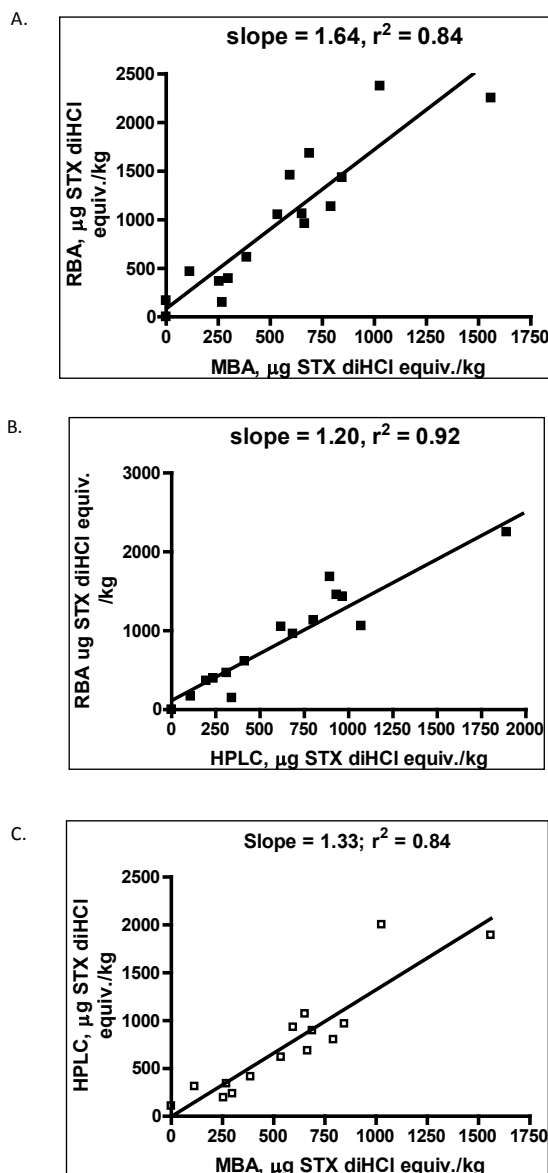


Figure 3. Correlation of the RBA results on PSP toxins in shellfish homogenates with mouse bioassay (A) and HPLC (B). Correlation between the current AOAC Official Methods, mouse bioassay, and HPLC (C).

the overall interlaboratory reproducibility can be further improved. It is recommended that this method be accepted by AOAC INTERNATIONAL as Official First Action for the determination of PST in shellfish.

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Appendix: Rat Brain Membrane Preparation

The rat brain membrane preparation used in this assay can be produced in bulk, aliquotted, and stored at -80°C until use. Under this storage condition, the preparation is stable for a minimum of 6 months. The following protocol provides sufficient membrane preparation for a minimum of 125 plates and can be scaled up or down as needed.

A. Apparatus

- (a) *Teflon/glass homogenizer*.—Motorized tapered Teflon pestle and glass tube, 15 mL.
- (b) *Motorized tissue homogenizer*.—Polytron or small hand-held blender.
- (c) *High-speed centrifuge and fixed angle rotor*.—Capable of $20\,000 \times g$ (rcf).
- (d) *Centrifuge tubes*.—12–15 mL rated for $>20\,000 \times g$ (rcf).
- (e) *Plastic cryovials*.—2 mL.
- (f) *Graduated beaker*.—300 or 500 mL.
- (g) *Pipets*.—Disposable 5 and 10 mL.
- (h) *Forceps*.

B. Reagents

- (a) *20 Rat brains*.—Male, 6-week-old Sprague-Dawley (Hilltop Lab Animals, Inc., Scottsdale, PA; <http://hilltoplabs.com>) or equivalent.
- (b) *MOPS*.—pH 7.4 (Sigma, St. Louis, MO; Cat. No. M3183-500G).
- (c) *Choline chloride*.—100 mM (Sigma; Cat. No. C7527-500G).
- (d) *Phenyl methylsulfonyl fluoride (PMSF)*.—Sigma; Cat. No. P7626.
- (e) *Isopropanol*.

C. Procedure

(1) Prepare 1 L 100 mM MOPS buffer, pH 7.4, containing 100 mM choline chloride (detailed protocol in E, above) and 0.1 mM PMSF. PMSF must first be dissolved in isopropanol; dissolve 0.174 g PMSF in 10 mL isopropanol to make 100 mM stock. Aliquot and store at -20°C . Add PMSF (1/1000, 0.1 mM final concentration) to the MOPS/choline chloride buffer fresh on the day of use.

(2) Remove medulla and cerebellum from each brain using forceps and discard. Place the cerebral cortex (see Figure 1) in a small amount of ice-cold buffer and place on ice.

(3) Place one cerebral cortex in 12.5 mL MOPS/choline Cl/PMSF, pH 7.4, in glass/teflon homogenizer (two brains in 25 mL buffer will fit into 30 mL homogenizer tube). Homogenize at 70% full speed (385 rpm) with at least 10 up and down strokes (more if necessary to homogenize brain; there should be no visible chunks remaining in the homogenate). Keep tube in ice at all times. Pour homogenized tissue into 250 mL beaker on ice and repeat procedure with remaining cortices.

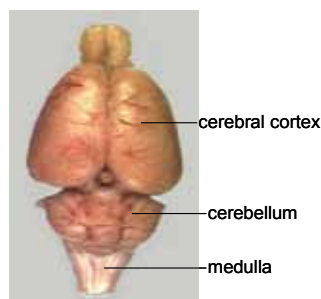


Figure 1. Rat brain.

(4) Transfer pooled homogenized tissue to centrifuge tubes, balance the tubes (pairwise; use ice-cold buffer to balance), and centrifuge at $20\,000 \times g$ for 15 min at 4°C .

(5) Aspirate the supernatant and resuspend the pellets in ice-cold MOPS/choline Cl/PMSF buffer, using an adequate amount (~ 5 mL) to fully resuspend the pellet (can use clean glass stir rod to break up pellet), not exceeding 10 mL per brain.

(6) Pool resuspended membrane preparation in a small beaker. Rinse centrifuge tubes with a small amount of ice-cold buffer to recover all of the membrane preparation. Bring total volume to 200 mL total (keep on ice).

(7) Keeping the beaker on ice, Polytron (or use a small hand-held blender at low speed) at 70% full speed for 20 s to obtain a consistent homogenate.

(8) Aliquot 2 mL/tube into cryovials. It is critical to keep the preparation well mixed while dispensing, e.g., prior to each aliquot to ensure equal allocation of protein/receptors to each vial. Keep cryotubes on ice.

(9) Freeze and store at -80°C . This preparation is stable for at least 6 months. Use a permanent marker to label the preparation date on the storage container.

D. Protein Assay

(a) Determine protein concentration of membrane preparation using Pierce Micro BCA Protein Assay Reagent Kit No. 23235 (microplate method) or No. 23225 (tube method) protein assay kit or equivalent protein assay (Thermo Fisher, Rockford, IL). The above protocol should yield 6–8 mg protein/mL of rat membrane preparation.

(b) Determine membrane dilution needed for the assay. The protein concentration in the daily working stock for the assay should be 1 mg/mL (this is diluted in the assay to yield 0.5 mg/mL in-assay concentration). Based on the protein concentration determined in the protein assay, determine the dilution needed to achieve 1 mg/mL. This is the dilution used in section E(e) above for all assays using this lot of membrane preparation. The protocol above typically yields a protein concentration that requires a dilution of 1/6–1/8. (Do not use less than 1/4 dilution or filtration wells may become clogged.) Protein concentration will need to be determined for each new batch of membrane preparation.

AOAC Official Method 2011.27
Paralytic Shellfish Toxins (PSTs) in Shellfish
Receptor Binding Assay
First Action 2011

[Applicable to the determination of paralytic shellfish toxins (PSTs), as μg STX diHCl equiv./kg, in shellfish (mussels, clams, scallops) at levels $>149 \mu\text{g}$ STX diHCl equiv./kg, with a limit of detection (LOD) of 45 STX diHCl equiv./kg shellfish and a limit of quantification (LOQ) of 126 μg STX diHCl equiv./kg shellfish.]

Caution: Wear disposable gloves and protective laboratory coat while performing the assay. PSTs are neurotoxins that are harmful if ingested. The assay uses a tritium labeled tracer, [^3H] STX, at low concentration. All laboratories performing the assay must have approved radiation laboratory space and must follow procedures prescribed by their nuclear regulatory agency for receipt, use, and disposal of isotopes.

See Tables 2011.27A–E for the results of the interlaboratory study supporting acceptance of the method.

A. Principle

Test portions of shellfish homogenates are extracted using the AOAC mouse bioassay extraction protocol (959.08), modified by scale. The PST receptor assay is a competitive binding assay in which [^3H] STX competes with unlabeled STX in standards or mixtures of PST in samples for a finite number of available receptor sites (site 1 on the voltage gated sodium channel) in a rat brain membrane preparation. Following establishment of binding equilibrium at 4°C, unbound [^3H] STX is removed by filtration and bound [^3H] STX is quantified by liquid scintillation counting. A standard curve is generated using increasing concentrations of STX standard from 10^{-10} to 10^{-6} M STX, which results in a reduction in bound [^3H] STX that is directly proportional to the amount of unlabeled toxin present. The concentration of toxin in samples is determined in reference to the standard curve. Incubation is carried out in a microplate format to minimize sample handling and the amount of radioactivity used. Bound [^3H] STX (as counts per minute; CPM) can be determined either by conventional or by microplate scintillation counting. Both methods are included in this protocol.

B. Apparatus and Supplies

- (a) *Traditional or microplate scintillation counter.*
- (b) *Micropipettors.*—1–1000 μL variable volumes and disposable tips.
- (c) *Eight channel pipettor.*—5–200 μL variable volume and disposable tips.
- (d) *96-Well microtiter filter plate.*—With 1.0 μm pore size type GF/B glass fiber filter/0.65 μm pore size Durapore support membrane (Millipore, Bedford, MA, USA; Cat. No. MSFB N6B 50).
- (e) *MultiScreen vacuum manifold.*—Millipore; Cat. No. NSVMHTS00.
- (f) *Vacuum pump.*
- (g) *Centrifuge tubes.*—15 and 50 mL, conical, plastic.
- (h) *Mini dilution tubes in 96-tube array.*
- (i) *Reagent reservoirs.*
- (j) *Ice bucket and ice.*
- (k) *Vortex mixer.*

- (l) *Sealing tape.*—Millipore; Cat. No. MATA HCL00.
- (m) *Volumetric flask.*—1 L.
- (n) *–80°C freezer.*
- (o) *Refrigerator.*

For traditional scintillation counter only:

- (p) *MultiScreen punch device.*—Millipore; Cat. No. MAMP 096 08.
 - (q) *MultiScreen disposable punch tips.*—Millipore; Cat. No. MADP 196 10.
 - (r) *MultiScreen punch kit B for 4 mL vials.*—Millipore; Cat. No. MAPK 896 0B.
 - (s) *Scintillation vials.*—4 mL.
- For sample extraction:
- (t) *Pipets.*
 - (u) *Centrifuge tubes.*—15 mL, conical, plastic.
 - (v) *Vacuum pump or house vacuum.*
 - (w) *pH meter or pH paper.*
 - (x) *Hot plate.*
 - (y) *Graduated centrifuge tubes.*—15 mL.
 - (z) *Centrifuge and rotor for 15 mL tubes.*

C. Reagents

- (a) [^3H] STX.—0.1 mCi/mL, ≥ 10 Ci/mmol, $\geq 90\%$ radiochemical purity (American Radiolabeled Chemicals, St. Louis, MO, USA, or International Isotopes Clearinghouse, Leawood, KS, USA).
 - (b) *STX diHCl.*—NIST RM 8642 (www.nist.gov).
 - (c) *3-Morpholinopropanesulfonic acid (MOPS).*—Sigma (St. Louis, MO, USA; Cat. No. M3183-500G), or equivalent.
 - (d) *Choline chloride.*—Sigma (Cat. No. C7527-500G), or equivalent.
 - (e) *Rat brain membrane preparation.*—Appendix 1 [*J. AOAC Int.* (future issue)].
- For traditional counter:
- (f) *Scintiverse BD liquid scintillation cocktail.*—Fisher Scientific (Waltham, MA, USA; Cat. No. SX-18), or equivalent.
- For microplate counter:
- (g) *Optiphase liquid scintillation cocktail.*—PerkinElmer Life Sciences (Downers Grove, IL, USA; Cat. No. 1200-139), or equivalent.
- For sample extraction:
- (h) *Hydrochloric acid (HCl).*—1.0 and 0.1 M.
 - (i) *Sodium hydroxide.*—0.1 M.
 - (j) *Water.*—Distilled or deionized (18 $\mu\Omega$).

D. Sample Extraction

Accurately weigh 5.0 g tissue homogenate into a tared 15 mL conical tube. Add 5.0 mL of 0.1 M HCl, vortex, and check pH. If necessary, adjust pH to 3.0–4.0 as determined by a pH meter or pH paper. To lower pH, add 1 M HCl dropwise with mixing; to raise pH, add 0.1 M NaOH dropwise with mixing to prevent local alkalization and consequent destruction of toxin. Place the tube in a beaker of boiling water on hot plate for 5 min with the caps loosened. Remove and cool to room temperature. Check pH and adjust cooled mixture to pH 3.0–4.0 as described above. Transfer entire contents to graduated centrifuge tube and dilute volumetrically to 10 mL. Gently stir contents to homogeneity and allow to settle until portion of supernatant is translucent and can be decanted free of solid particles. Pour approximately 5 to 7 mL of the translucent supernatant into a centrifuge tube. Centrifuge at $3000 \times g$ for 10 min. Retain clarified supernatant and transfer to a clean centrifuge tube. Store extracts at -20°C until tested in

Table 2011.27A. Receptor binding assay results on individual samples (values are in µg STX diHCl equiv./kg shellfish tissue); summary statistics on all samples; summary statistics excluding Laboratory 9

Assay	No.	ID	Sample									All labs					Labs 1-8		
			1	2	3	4	5	6	7	8	9	Mean	S _R	RSD _R , %	HorRat	Mean	S _R	RSD _R , %	HorRat
Day 1	1	MLV05	370	610	620	410	690	1070	630	660	330	599	222	37.1	2.2	633	212	33.5	2.0
	2	MLV06	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	422	28.6	1.9	1549	386	24.9	1.7
Day 2	3	MLV08	80	190	140	90	130	160	230	220	100	149	55	37.2	1.8	155	56	36.0	1.7
	4	MLV09	860	680	950	870	980	1120	1460	820	590	926	255	27.5	1.7	968	237	24.5	1.5
Day 3	5	MLV12	180 ^a	200	200	150	150	100	150	290	100	168	62	37.2	1.8	177	60	34.1	1.7
	6	MLV14	950	940	1060	1130	1040	750	1460	1320	810	1051	228	21.7	1.4	1081	224	20.7	1.3
Day 3	7	MLV16	660	930	1080	870	840	1320	1490	2420 ^b	490	960	329	34.3	2.1	1027	291	28.3	1.8
	8	MLV01	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	818	48.6	3.3	1829	739	40.4	2.8
Day 3	9	MVL02	830	1180	1130	1150	1130	1780	1340	980	690	1134	311	27.4	1.8	1190	281	23.6	1.5
	10	MLV04	2440	2840	2910	1740	2150	1810	2690	2490	1210	2253	572	25.4	1.8	2384	446	18.7	1.3
Day 3	11	MLV07	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	345	23.9	1.6	1520	279	18.3	1.2
	12	MLV09	810	1190	1130	810	1630	1390	1880	1120	870	1203	372	30.9	2.0	1245	375	30.1	2.0
Day 3	13	MLV11	270	370	480	340	640	490	240	600	110	393	174	44.3	2.4	429	148	34.4	1.9
	14	MLV13	400	1240 ^b	560	450	650	530	500	440	200	466	133	28.5	1.6	504	85	16.8	1.0
Day 3	15	MLV03	330	270	410	180	590	680	370	1570 ^b	90	365	197	54.0	2.9	404	176	43.5	2.4
	16	MLV05	580	670	250	430	910	700	860	940	300	627	257	41.1	2.4	668	242	36.2	2.1
Day 3	17	MLV06	1290	1520	1460	970	1800	2520	1470	870	1250	1461	488	33.4	2.2	1488	515	34.6	2.3
	18	MLV07	1010	1600	1390	1000	1720	1860	1520	2150	890	1460	429	29.4	2.0	1531	397	26.0	1.7
Day 3	19	MLV10	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	570	24.0	1.7	2443	569	23.3	1.7
	20	MLV11	430	350	460	280	550	620	1149 ^b	410	250	419	127	30.2	1.7	443	115	26.0	1.4
Day 3	21	MLV15	ND ^c	ND	ND	ND	ND	ND	ND	180	ND	—	—	—	—	—	—	—	—
	Avg. RSD _R																		
	Avg. HorRat												33.2	2.0					28.7

^a CV 41%, not used in calculations.

^b Outlier; not used in calculations.

^c ND = Not detected.

Table 2011.27B. Summary statistics on blind duplicates, run in separate assays (values are in µg STX diHCl equiv./kg)

Lab	MLV05		MLV06		MLV07		MLV09		MLV11		Avg.
	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	
1	370	580	1100	1290	1260	1010	860	810	270	430	
2	610	670	1340	1520	1540	1530	680	1190	370	350	
3	620	250	1320	1460	1220	1390	950	1130	480	401	
4	410	430	1440	970	1980	1000	870	810	340	280	
5	690	910	1260	1790	1760	1720	980	1630	640	550	
6	1070	700	1720	2520	1530	1860	1120	1390	490	620	
7	630	880	2090	1240	1750	1150	1460	1830	230 ^a	1149 ^a	
8	660	940	2130	870	1210	2150	820	1120	600	410	
9	330	300	890	1250	840	890	590	870	110	250	
Avg.		614		1453		1433		1062		416	
S _r		169		432		366		247		83	
S _R		239		444		387		338		152	
RSD _r %		27.5		29.4		25.5		23.3		20.0	25.1
RSD _R %		38.9		30.2		27.0		31.9		36.5	32.9
HorRat		2.3		2.0		1.8		2.0		2.0	2.0

^a Outlier; not used in calculation.

receptor assay.

E. Preparation of Stock Solutions and Standards

(a) *Assay buffer*.—100 mM MOPS/100 mM choline chloride, pH 7.4. Weigh out 20.9 g MOPS and 13.96 g choline chloride and add to 900 mL dH₂O. Adjust pH to 7.4 with NaOH while stirring and bring to a final volume of 1 L with dH₂O. Store at 4°C.

(b) *Radioligand solution*.—Calculate the concentration of [³H] STX stock provided by the supplier, which may vary between lots. Suppliers generally provide the specific activity in Ci/mmol (generally 10–30 Ci/mmol) and activity in mCi/mL (0.05–0.1 mCi/mL), from which the molar concentration can be calculated. Prepare 4 mL of a 15 nM working stock of [³H] STX fresh daily in 100 mM MOPS/100 mM choline chloride buffer. This will provide sufficient volume for one 96-well plate at an in-well concentration of 2.5 nM. Measure total counts of each working stock prior to running an assay: add 35 µL of the working stock [³H] STX in buffer to a liquid scintillation vial with 4 mL scintillant and count on a traditional liquid scintillation counter. This is done to confirm correct dilution prior to running the assay. Depending on the efficiency of the scintillation counter used, the corresponding CPM will vary, but should be consistent day-to-day and within 15% of the expected value.

(c) *Unlabeled STX standard working solution*.—The STX diHCl standard is provided at a concentration of 268.8 µM (100 µg/mL). A “bulk” standard curve can be made up in advance and stored at 4°C for up to 1 month. The use of a bulk standard curve minimizes the pipetting needed for setting up an assay routinely and improves day-to-day repeatability. Make up 3 mM HCl (e.g., from a 3 M stock, 50 µL in 50 mL), then perform the serial dilutions (see Table 2011.27F) of NIST RM 8642 STX diHCl (100 µg/mL = 268.8 µM) to make up the standard curve in 3 mM HCl. These standard stock solutions will be diluted 1/6 in the assay to yield the designated in-assay concentrations (see Table 2011.27F).

(d) *Interassay calibration standard (QC check)*.—Prepare a

reference standard containing 1.8×10^{-8} M STX standard (3.0×10^{-9} M STX in assay) in advance in 3 mM HCl and keep frozen (–80°C) in 1 mL aliquots for long-term storage. Aliquots should be thawed and stored at 4°C for routine use (stable up to 1 month) and analyzed in each assay. This serves as a QC check and confirms day-to-day performance of the assay.

(e) *Rat brain membrane preparation*.—Prepare rat brain membrane preparation in bulk [Appendix I; J. AOAC Int. (future issue)] and store at –80°C until used in the assay. Thaw an aliquot of rat brain membrane preparation on ice. Dilute membrane preparation with cold (4°C) 100 mM MOPS/100 mM choline chloride, pH 7.4, to yield a working stock with a protein concentration of 1.0 mg/mL (this will be diluted in the assay plate to 0.5 mg/mL in-well concentration). Vortex vigorously to achieve a visibly homogeneous suspension. Keep the diluted membrane preparation on ice until ready to use.

F. Performing the Assay

(a) *Plate setup*.—When possible, use a multichannel pipet to minimize pipetting effort and increase consistency. Standard curve, QC check, and sample extracts are run in triplicate wells. Multiple dilutions of sample extracts should be analyzed in order to obtain a value that falls between 0.2–0.7 B/B₀ on the standard curve for quantification. For ease of analysis, it is convenient to use a standard plate layout that maximizes the number of samples and standards that can be analyzed on one plate. For shellfish extracts, a minimum dilution of 1:10 is used, which minimizes potential matrix effects, while still providing an LOQ of approximately 126 µg/kg shellfish (see Table 2011.27G).

(b) *Addition of samples and standards*.—Add in the following order to each of the 96 wells: 35 µL assay buffer; 35 µL STX standard, QC check, or sample extract; 35 µL [³H] STX; 105 µL membrane preparation. The assay buffer is added first in order to wet the filter membrane. It is critical to continuously mix the membrane preparation by careful up-and-down pipetting immediately prior to

Table 2011.27C. Performance of individual laboratories on blind duplicates (values are in µg STX diHCl equiv./kg)

Laboratory	ID	Day 1	Day 2	Mean	s _p	RSD _r , %
1	MLV05	370	580	475	148	31.3
	MLV06	1100	1290	1195	134	11.2
	MLV07	1260	1010	1135	177	15.6
	MLV09	860	810	835	35	4.2
	MLV11	270	430	350	113	32.3
Avg.						18.9
2	MLV05	605	670	638	46	7.2
	MLV06	1340	1520	1430	127	8.9
	MLV07	1540	1530	1535	7	0.5
	MLV09	680	1190	935	361	38.6
	MLV11	370	350	360	14	3.9
Avg.						11.8
3	MLV05	620	250	435	262	60.1
	MLV06	1320	1460	1390	99	7.1
	MLV07	1220	1303	1262	59	4.7
	MLV09	950	1130	1040	127	12.2
	MLV11	480	460	470	14	3.0
Avg.						17.4
4	MLV05	410	430	420	14	3.4
	MLV06	1440	970	1205	332	27.6
	MLV07	1980	1000	1490	693	46.5
	MLV09	870	810	840	42	5.1
	MLV11	340	280	310	42	13.7
Avg.						19.2
5	MLV05	690	910	800	156	19.4
	MLV06	1260	1790	1525	375	24.6
	MLV07	1760	1720	1740	28	1.6
	MLV09	980	1630	1305	460	35.2
	MLV11	640	550	595	64	10.7
Avg.						18.3
6	MLV05	1070	700	885	262	29.6
	MLV06	1720	2520	2120	566	26.7
	MLV07	1530	1860	1695	233	13.8
	MLV09	1120	1390	1255	191	15.2
	MLV11	490	620	555	92	16.6
Avg.						20.4
7	MLV05	630	880	755	177	23.4
	MLV06	2090	1240	1665	601	36.1
	MLV07	1750	1150	1450	424	29.3
	MLV09	1460	1830	1645	262	15.9
	MLV11	230 ^a	1150 ^a			
Avg.						26.2
8	MLV05	660	940	800	198	24.7
	MLV06	2130	870	1500	891	59.4
	MLV07	1210	2150	1680	665	39.6
	MLV09	820	1120	970	212	21.9
	MLV11	600	410	505	134	26.6
Avg.						34.4
9	MLV05	330	300	315	21	6.7
	MLV06	890	1250	1070	255	23.8
	MLV07	840	890	865	35	4.1
	MLV09	590	870	730	198	27.1
	MLV11	110	250	180	99	55.0
Avg.						23.3
Overall avg.						22.2

^a Outlier; not used in calculations.

dispensing into the 96-well plate to maintain an even suspension across the entire plate. Cover and incubate plate at 4°C for 1 h.

(c) *Assay filtration.*—Attach the vacuum manifold to the vacuum pump with an in-line side arm flask to catch filtrate from the plate filtration process. Set the vacuum pressure gauge on the pump or vacuum manifold to 4–8" Hg (135–270 millibar), as specified in the instructions provided with the filtration plates. Place the 96-well plate on the vacuum manifold. Fill empty wells with 200 µL MOPS/choline chloride buffer to ensure even vacuum pressure and filtration across the plate. Turn on vacuum. Optimum vacuum will pull the wells to dryness in 2–5 s. Pull contents of all wells through until all liquid is removed. (*Note:* Too low a vacuum will result in slow well clearance, but too high will result in an airlock and no well clearance.) With vacuum pump running, quickly rinse each well twice with 200 µL ice cold MOPS/choline chloride buffer using multichannel pipet. Maintain vacuum until liquid is removed.

(d) *Preparation of the assay for counting.*—Remove the plastic bottom from the plate. Blot the bottom once on absorbent toweling.

(1) *For counting in microplate scintillation counter.*—Place the microplate in a counting cassette. Seal the bottom of the 96-well plate with sealing tape. Add 50 µL Optiphase scintillation cocktail per well using multichannel pipet. Seal the top of the plate with sealing tape. Allow to incubate 30 min at room temperature. Place the plate in a counting cassette and count in a microplate scintillation counter for 1 min/well.

(2) *For counting in traditional scintillation counter.*—Place the microplate in the MultiScreen punch system apparatus. Place the disposable punch tips on top of the microplate. Punch the filters from the wells into scintillation vials and fill with 4 mL scintillation cocktail (Scintiverse or equivalent). Place caps on the vials and vortex. Allow vials to sit overnight in the dark, then count using a tritium window in a traditional scintillation counter.

G. Analysis of Data

For assays performed using the traditional counter, curve fitting is performed using a four-parameter logistic fit, also known as a sigmoidal dose response curve (variable slope; *see* Figure 2011.27), or Hill equation:

$$y = \min + \frac{\max - \min}{1 + 10^{(x - \log IC_{50}) \text{Hill slope}}}$$

where max is the top plateau representing maximum binding in CPM in the absence of competing nonradiolabeled STX, also known as B₀; min is the bottom plateau, equal to nonspecific binding (in CPM) in the presence of saturating nonradiolabeled toxin; IC₅₀ is the inhibitory concentration at which CPM are 50% of max-min (dashed lines; Figure 2011.27); Hill slope is the slope of the curve; x axis is the log concentration of STX; and y axis is total ligand binding in CPM (here represented as B/B₀, or bound/max bound). A curve fitting package such as Prism (Graph Pad Software, Inc.) is recommended. For the microplate counter users, receptor assay applications provided by the manufacturer may be used (e.g., MultiCalc; PerkinElmer Wallac, Gaithersburg, MD, USA).

(a) *Sample quantification.*—Sample quantification is carried out only on dilutions that fall within B/B₀ of 0.2–0.7, where B represents the bound [³H]STX (in CPM) in the sample and B₀ represents the max bound [³H]STX (in CPM). Where more than one dilution falls within B/B₀ of 0.2–0.7 on the curve, all sample wells corresponding to these dilutions are used to calculate sample concentration. Sample concentration is calculated in µg STX diHCl

Table 2011.27D. Calibration curve and QC check parameters in three receptor binding assays performed in nine participant laboratories

Lab	Assay day	Slope	IC ₅₀ ^a , nM	QC, nM	Reference, CPM	IC ₇₀ ^a , nM	Standards where RSD >30%; action	Curve fitting software	Scintillation counter	Manual/microplate
1	1	-0.9	1.9	2.4	720	0.90	None	Prism v 3.02	Packard Top Count	Microplate
	2	-1.0	2.0	2.6	733	0.96	None			
	3	-1.1	2.1	3.2	1038	0.92	None			
2	1	-1.1	1.8	3.8	1160	0.66	3 nM; 1 well removed	Prism v 5.0	Packard Top Count	Microplate
	2	-1.2	2.2	3.9	1260	0.85	None			
	3	-1.0	1.6	3.2	1262	0.46	3 nM, 1 nM removed			
3	1	-1.0	2.0	2.3	2529	0.41	First column removed	Prism v 5.0	Wallac Microbeta	Microplate
	2	-0.9	2.0	2.5	1463	0.92	1000 nM; 1 well removed			
	3	1.0	1.6	2.8	2088	0.80	None			
4	1	-0.9	1.7	3.4	1125	0.61	None	Prism v 3.03	PerkinElmer Tricarb	Manual
	2	-1.2	1.7	3.2 ^a	1611	0.77	None			
	3	-0.9	1.2	2.9	1324	0.45	30 nM 35%; 1 well removed			
5	1	-0.9	1.4	3.3	1566	0.64	1.0 nM; 1 well removed	MultiCalc	Wallac Microbeta	Microplate
	2	-1.2	1.8	3.6	1528	1.05	0.1 nM and 30 nM; 1 well removed			
	3	-1.2	1.8	2.9	1052	0.67	None			
6	1	-1.1	2.6	3.0	670	1.15	None	Prism v 4.0	Wallac Microbeta	Microplate
	2	-1.0	2.0	4.0 ^b	1124	1.08	None			
	3	-1.1	3.4	6.5 ^b	1030	2.04 ^c	None			
7	1	-0.8	1.0	2.8 ^a	919	0.33	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	1.6	2.7	619	0.70	None			
	3	-0.9	2.1	3.2 ^a	693	0.82	None			
8	1	-1.2	1.7	3.7	1146	0.86	None	Prism	Wallac Microbeta	Microplate
	2	-1.1	1.4	1.5 ^b	1095	0.78	None			
	3	-1.1	2.4	2.3	886	1.04	None			
9	1	-1.0	2.2	4.0 ^b	1363	0.97	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	2.0	3.2	1380	0.85	100 nM 33%; left in			
	3	-1.0	2.1	3.7	1532	0.92	None			

^a One well removed.

^b Outside of specifications.

^c Outlier by Grubbs test.

Table 2011.27E. Results of the receptor binding assay (RBA), mouse bioassay (MBA), and HPLC analyses of 21 shellfish extracts, sorted by mouse bioassay value (all values are in µg STX diHCl equiv./kg shellfish tissue; results in bold indicate toxicity above the 800 µg STX diHCl equiv./kg regulatory limit; all other results indicate toxicity below the regulatory limit)

Sample	Lab 1	Lab 2	Lab 3	Lab 4	Lab 5	Lab 6	Lab 7	Lab 8	Lab 9	RBA, avg.	HPLC	MBA
21	ND ^a	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
5	180	200	200	150	150	100	150	290	100	168	108	ND
15	330	270	410	180	590	680	370	1570^b	90	365	196	182
13	270	370	480	340	640	290	240	600	110	371	236	299
20	430	350	460	280	550	490	1150^b	410	250	403	236	299
14	400	1240^b	560	450	650	530	500	440	200	466	625	343
1	370	610	620	410	690	1070^b	630^b	660	330	599	413	387
16	580	670	250	430	910	700	860^b	940^b	300	627	413	387
3	80	190	140	90	130	160	230	220	100	149	341	405
6	950	940	1060	1130	1040	750	1460	1320	810	1051	618	485
7	660	930	1080	870	840	1320	1490	2420	490	960	685	528
2	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	931	595
17	1290	1520	1460	970	1800	2520	1470	870	1250	1460	931	595
4	860	680	950	870	980	1120	1460	820	590	926	1070	653
12	810	1190	1130	810	1630	1390	1880	1120	870	1203	1070	653
11	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	965	714
18	1010	1600	1390	1000	1720	1860	1520	2150	890	1452	965	714
8	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	894	752
9	830	1180	1130	1150	1130	1780	1340	980	690	1134	802	792
19	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	2000	1027
10	2440	2840	2910	1740	2150	1800	2690	2490	1210	2252	1890	1080

^a ND = Not detected.

^b Outlier; not used in average calculation.

equiv./kg shellfish, using the following formulas:

$$(\text{nM STX equiv.}) \times (\text{sample dilution}) \times \frac{(210 \mu\text{L total volume})}{35 \mu\text{L sample}} = \text{nM STX equiv. in extract}$$

$$(\text{nM STX diHCl equiv. in extract}) \times \frac{1 \text{ L}}{1000 \text{ mL}} \times \frac{372 \text{ ng}}{\text{nmol}} \times \frac{1 \mu\text{g}}{1000 \text{ ng}} = \mu\text{g STX diHCl equiv./mL}$$

$$\mu\text{g STX diHCl equiv./mL} \times \frac{\text{mL extract}}{\text{g shellfish}} \times \frac{1000 \text{ g}}{\text{kg}} = \mu\text{g STX diHCl equiv./kg}$$

H. Assay Performance Standards

The following criteria must be met for assay acceptance:

(a) For a ligand that specifically binds at one receptor site, the slope of the resulting competition curve should theoretically be -1.0. If the slope of the curve for a given assay is outside of the acceptable range of -0.8 to -1.2, linearity of the assay will be compromised and quantification of the unknowns will be incorrect.

(b) RSDs of triplicate CPMs for standards should be below 30% as variability may affect the slope calculation and thereby quantification of samples.

(c) If the IC₅₀ is out of the acceptable range (2.0 nM ± 30%)

then the assay should be considered suspect and rerun, as a shift in the curve will result in over- or underestimation of sample concentrations.

(d) QC check should be 3 nM STX ± 30% (in-well concentration).

Table 2011.27F. Dilution series to prepare bulk solutions for standard curve

	Stock, M	In-assay, M
100 µL 268.8 µM STX + 4.38 mL 0.003 M HCl	6 × 10 ⁻⁶	1 × 10 ⁻⁶
500 µL 6 × 10 ⁻⁶ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁷	1 × 10 ⁻⁷
1.5 mL 6 × 10 ⁻⁷ M + 3.5 mL 0.003 M HCl	1.8 × 10 ⁻⁷	3 × 10 ⁻⁸
500 µL 6 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁸	1 × 10 ⁻⁸
500 µL 1.8 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	1.8 × 10 ⁻⁸	3 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁸ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁹	1 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁹ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻¹⁰	1 × 10 ⁻¹⁰
5 mL 0.003 M HCl	0	Reference

Table 2011.27G. Recommended microplate layout for ease of handling triplicate wells of standard curve, QC check sample, and unknown samples; each sample is run at three dilutions (1:10, 1:50, 1:200); standard curve is run in columns 1–3 (values are in M STX)^a

Microplate row	Microplate column											
	1	2	3	4	5	6	7	8	9	10	11	12
A	10 ⁻⁶	10 ⁻⁶	10 ⁻⁶	QC	QC	QC	U3 1:50	U3 1:50	U3 1:50	U6 1:10	U6 1:10	U6 1:10
B	10 ⁻⁷	10 ⁻⁷	10 ⁻⁷	U1 1:10	U1 1:10	U1 1:10	U3 1:200	U3 1:200	U3 1:200	U6 1:50	U6 1:50	U6 1:50
C	3 × 10 ⁻⁸	3 × 10 ⁻⁸	3 × 10 ⁻⁸	U1 1:50	U1 1:50	U1 1:50	U4 1:10	U4 1:10	U4 1:10	U6 1:200	U6 1:200	U6 1:200
D	10 ⁻⁸	10 ⁻⁸	10 ⁻⁸	U1 1:200	U1 1:200	U1 1:200	U4 1:50	U4 1:50	U4 1:50	U7 1:10	U7 1:10	U7 1:10
E	3 × 10 ⁻⁹	3 × 10 ⁻⁹	3 × 10 ⁻⁹	U2 1:10	U2 1:10	U2 1:10	U4 1:200	U4 1:200	U 1:200	U7 1:50	U7 1:50	U7 1:50
F	10 ⁻⁹	10 ⁻⁹	10 ⁻⁹	U2 1:50	U2 1:50	U2 1:50	U5 1:10	U5 1:10	U5 1:10	U7 1:200	U7 1:200	U7 1:200
G	10 ⁻¹⁰	10 ⁻¹⁰	10 ⁻¹⁰	U2 1:200	U2 1:200	U2 1:200	U5 1:50	U5 1:50	U5 1:50			
H	REF	REF	REF	U3 1:10	U3 1:10	U3 1:10	U5 1:200	U5 1:200	U5 1:200			

^a REF = Reference; QC = quality control check; U = unknown sample. [Note: The same standard curve may be used for multiple plates (i.e., 11 samples can be run on subsequent plates in a series if the standard curve is not included).]

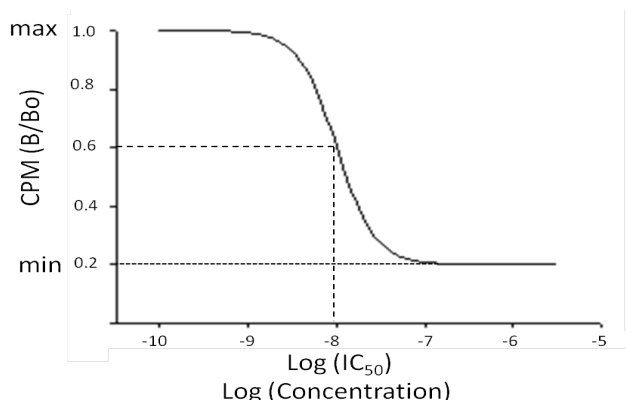


Figure 2011.27. Sigmoidal dose response curve. Dashed lines indicate log IC50.

Assays with a QC check sample out of specifications should trigger a check of the IC₅₀ value.

The following criteria must be met for acceptability of a sample measurement:

(a) Sample quantification should be done only on dilutions that fall within B/B₀ of 0.2–0.7. In the event that all sample dilutions fall below B/B₀ 0.2 (i.e., concentration is too high), further dilutions must be made and the sample reanalyzed. In the event that the sample concentration is too low to be quantified (i.e., B/B₀ > 0.7), the sample is reported as below LOD. If more than one dilution falls on the linear part of the curve, an average value calculated from all dilutions should be used. If there is disagreement between different dilutions in final concentration reported, check for error in the sample dilution process.

(b) RSD of the sample CPMs should be ≤30%.

Reference: *J. AOAC Int.* (future issue)



Report of Investigation

Reference Material 8642

FDA Saxitoxin Dihydrochloride Solution

This Reference Material (RM) is intended for use in calibrating the mouse bioassay used in AOAC International Official Method 959.08 Paralytical Shellfish Poison [1] and for other similar uses. RM 8642 FDA Saxitoxin Dihydrochloride Solution was prepared by the U.S. Food and Drug Administration's (FDA's) Center for Food Safety and Applied Nutrition (CFSAN), where it was identified as Lot 089. The RM is saxitoxin dihydrochloride (CAS No. 35554-08-6) in a solution containing a hydrochloric acid concentration of 5 mmol/L in 20 % ethanol in water (volume fraction). A unit of RM 8642 consists of ten amber, borosilicate glass ampoules, each containing approximately 1.2 mL of solution.

Reference Mass Fraction Value: The reference value for the mass fraction of saxitoxin hydrochloride in solution in RM 8642, identified by FDA as lot 089, is 103 $\mu\text{g/g}$ with an expanded uncertainty of 4 $\mu\text{g/g}$. Reference values are noncertified values that are estimates of the true value; however, the values do not meet the NIST criteria for certification and are provided with associated uncertainties that may reflect only measurement precision, may not include all sources of uncertainty, or may reflect a lack of sufficient statistical agreement among multiple analytical methods [2]. The reference mass fraction value is based on the gravimetric preparation of a stock solution and gravimetric dilution to produce the final material, and uncertainties associated with the associated weighings. The uncertainty is expressed as an expanded uncertainty, $U = ku_c$, at the 95 % level of confidence, $k = 2$, and includes a 2 % Type B purity uncertainty component as well as the gravimetric uncertainty [3]. Values are reported on an "as-received" basis in mass fraction units [4].

Expiration of Value Assignment: The reference value for RM 8642 is valid, within the measurement uncertainty specified, until **01 July 2013**, provided the RM is handled and stored in accordance with instructions given in this report (see "Instructions for Use"). This report is nullified if the RM is damaged, contaminated, or otherwise modified.

Maintenance of RM: NIST will monitor this RM over the period of its validity. If substantive technical changes occur that affect the value assignment before the expiration of this report, NIST will notify the purchaser. Registration (see attached sheet) will facilitate notification.

The technical and support aspects involved in the preparation and issuance of this Reference Material were coordinated through K.E. Sharpless of the NIST Analytical Chemistry Division and M.P. Cronise of the NIST Measurement Services Division.

The solution was prepared and characterized by S. Hall of the Division of Bioanalytical Chemistry, Office of Regulatory Science, CFSAN, FDA.

Statistical analysis was provided by J.H. Yen of the NIST Statistical Engineering Division.

Support aspects involved in the issuance of this SRM were coordinated through the NIST Measurement Services Division.

Stephen A. Wise, Chief
Analytical Chemistry Division

Robert L. Watters, Jr., Chief
Measurement Services Division

Gaithersburg, MD 20899
Report Issue Date: 09 December 2010
Report Revision History on last page.

NOTICE AND WARNING TO USERS

Warning: For laboratory use only.

Storage: Unopened ampoules should be stored upright under normal laboratory conditions inside the original container supplied by NIST.

INSTRUCTIONS FOR USE

Gently tap the ampoule prior to opening to allow any solution in the tip to drain into the body of the ampoule.

Prepare a working solution as follows: On a top-loading balance, record the tare weight of an appropriate plastic bottle to 0.1 g or better. To the bottle, add approximately 100 mL water that has been acidified to pH 3 with hydrochloric acid. To minimize error due to evaporation, be prepared to immediately transfer the RM solution to this bottle after opening the ampoule. To open, hold the ampoule steady and grasp the stem at the metallic band with thumb and forefinger; **minimal** thumb pressure should be applied to the stem to snap it. Correctly done, the stem should break easily where pre-scored. Aspirate the RM solution into a dry, clean, disposable plastic syringe, 2 mL to 5 mL capacity, fitted with a suitable needle (such as 18 G \times 1 1/2"), weigh the syringe and its contents to 1 mg or better, and dispense the solution into the bottle of acidified water. Do not rinse the syringe. Reweigh the emptied syringe to determine the mass of RM solution transferred to the bottle. Add sufficient acidified water (pH 3, HCl) to adjust the concentration to 1 μ g/g. Weigh the bottle and its contents to determine the mass of solution prepared and the exact concentration of the working solution.

Because of the volatility of ethanol, the reference value is not applicable to material in ampoules that have been previously opened. The concentration of the working solution should be stable for more than one month if the solution is protected from evaporation. Dilution by mass is preferred but, if dilution by volume must be performed, the density of the solution is 0.971 g/mL and the concentration of this standard is 100 μ g/mL with an expanded uncertainty of 4 μ g/mL. This uncertainty is calculated as described above.

Source and Preparation of Material: Saxitoxin was extensively purified on three low-pressure preparative columns, each containing a different stationary phase. The saxitoxin was converted to the dihydrochloride form by passage through an ion exchange resin in the chloride form. Purity was assessed at FDA by proton nuclear magnetic resonance spectroscopy, combustion analysis, and optical rotation. RM 8642, identified by FDA as lot 089, was prepared by dissolving the saxitoxin dihydrochloride in a solution of hydrochloric acid (5 mmol/L) in 20 % ethanol in water (volume fraction).

REFERENCES

- [1] AOAC International; *Official Methods of Analysis of AOAC International*, 18th Edition, Gaithersburg, MD (2005).
- [2] May, W.; Parris, R.; Beck II, C.; Fassett, J.; Greenberg, R.; Guenther, F.; Kramer, G.; Wise, S.; Gills, T.; Colbert, J.; Gettings, R.; MacDonald, B.; *Definition of Terms and Modes Used at NIST for Value-Assignment of Reference Materials for Chemical Measurements*; NIST Special Publication 260-136 (2000); available at <http://ts.nist.gov/MeasurementServices/ReferenceMaterials/PUBLICATIONS.cfm> (accessed Nov 2010).
- [3] JCGM 100:2008; *Evaluation of Measurement Data — Guide to the Expression of Uncertainty in Measurement* (ISO GUM 1995 with Minor Corrections); Joint Committee for Guides in Metrology (2008); available at http://www.bipm.org/utis/common/documents/jcgm/JCGM_100_2008_E.pdf (accessed Nov 2010); see also Taylor, B.N.; Kuyatt, C.E.; *Guidelines for Evaluating and Expressing the Uncertainty of NIST Measurement Results*; NIST Technical Note 1297; U.S. Government Printing Office: Washington, DC (1994); available at <http://www.nist.gov/physlab/pubs/index.cfm> (accessed Nov 2010).
- [4] Thompson, A.; Taylor, B.N.; *Guide for the Use of the International System of Units (SI)*; NIST Special Publication 811; U.S. Government Printing Office: Washington, DC (2008); available at: http://ts.nist.gov/WeightsAndMeasures/Metric/mpo_pubs.cfm (accessed Nov 2010).

Report Revision History: 09 December 2010 (Extension of the period of validity; editorial changes.); 09 June 2010 (Original report date).
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Users of this RM should ensure that the Report of Investigation in their possession is current. This can be accomplished by contacting the SRM Program: telephone (301) 975-2200; fax (301) 926-4751; e-mail srminfo@nist.gov; or via the Internet at <http://www.nist.gov/srm>.

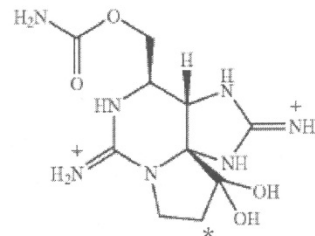


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 E-mail: arcinc@arc-inc.com

TECHNICAL DATA SHEET ART 1301 Saxitoxin [¹¹-³H]

LOT SPECIFIC TECHNICAL DATA:

Lot number: 120814
 Specific activity: estimated 20-30 Ci/mmol
 Solvent: Methanol
 Radioactive concentration: 0.05 mCi/ml
 Molecular weight: 299.2



PACKAGING INFORMATION:

ART 1301 is packaged as a solution in methanol in a sealed ampoule. It is shipped in dry ice.

STABILITY AND STORAGE RECOMMENDATIONS:

A working stock of 1/50 dilution in methanol can be stored at 4°C. Long-term storage should be carried out at -80°C, based on the previous commercially available Saxitoxin [³H], which was not stable at -20°C. The rate of degradation at -80°C is approximately 0.3-1% for the first month.

RADIOCHEMICAL AND CHEMICAL PURITY:

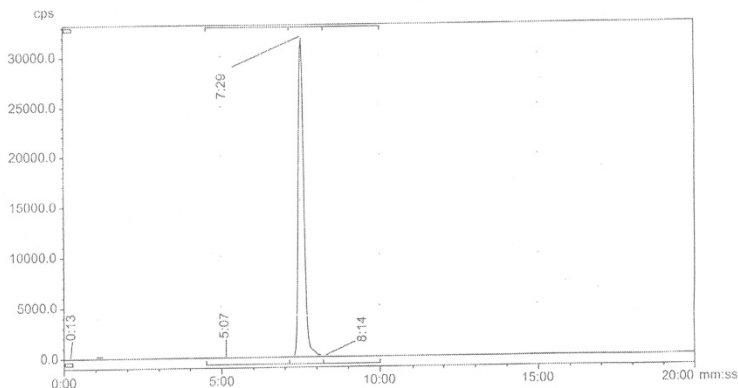
Radiochemical Purity: 99.56%

Column: Zorbax SB-AQ (250 x 3.0mm)

Mobile Phase: water:isopropanol:
 heptafluorobutyric acid
 (99.5 : 0.5 : 0.005)

Flow Rate: 0.5 ml/min

Detector: β-RAM [³H]



Name	Start (mm:ss)	End (mm:ss)	Retention (mm:ss)	Height (cps)	Area (Counts)	%ROI (%)
Bkg 1	0:02	0:16	0:13	183.0		
Region 1	4:31	7:08	5:07	158.0	1375.5	0.36
Region 2	7:08	8:11	7:29	32767.0	384313.4	99.56
Region 3	8:11	9:58	8:14	136.0	320.3	0.08
3 Peaks					386009.1	100.00

At the time of shipment all products are guaranteed to be free from defects in material and workmanship and to conform to the accompanying technical specifications and purity data. ARC will offer a 30 day money back guarantee of free replacement of products that are found to be unsatisfactory in respect to product specifications and purity. ARC makes no other warranty, expressed or implied, pertaining to the suitability of the product for any specific application. In case of breach of this warranty the entire liability of ARC will be limited to the invoice price of the goods. In no case will ARC be liable for any special, incidental or consequential damages resulting from the use of its products. ARC hereby expressly disclaims any warranty regarding results obtained through use of the products, including without limitation any claim of inaccurate, invalid, or incomplete results. Products are not suitable for human use.

Validation Data for Direct plating method for trh

Name of Method Submitter: Jessica L. Jones, Ph.D.

Specific purpose or intent of the method for use in the NSSP: Seeking approval for this method as an approved limited use method that can be used as appropriate for PHP validation and verification testing, as well as environmental testing such as that which may be required for the re-opening of growing areas closed due to illness.

Validation Criteria Data: For evaluation of all validation criteria below, PHP oysters were obtained in the best effort to find samples free of the target organism. A different lot of PHP oysters was used for each sample. For each sample, a minimum of 10 animals were used to prepare a homogenate. The homogenate was then aliquoted and appropriate aliquots spiked with a *tdh+/trh+ Vibrio parahaemolyticus* (unless otherwise noted), while one aliquot was left un inoculated (sample blank). Spike levels were determined by spread plating dilution of the culture in triplicate onto TSA+2% NaCl. Appropriate aliquots of spiked samples were spread plated onto T1N3 agar and colony lifts hybridized with an alkaline phosphatase-labeled probe specific for *trh*.

1. Accuracy/Trueness: Using the data from Table 1, the average of plate counts was 3.80 log and the average from DNA probe was 3.62 log. The Accuracy/Trueness of the method is 95%.

Table 1. Data for determination of Accuracy/Trueness and Measurement Uncertainty.		
Sample	Plate Count (log CFU)	Probe Result (log CFU/g)
1-2X	5.18	4.76
2-2X	5.18	4.65
3-4X	3.15	2.90
4-4X	3.15	2.85
5-6X	1.23	1.48
6-6X	1.23	1.00
7-2X	5.76	5.59
8-2X	5.76	5.64
9-4X	3.68	3.59
10-4X	3.68	3.72

2. Measurement Uncertainty: Using the data from Table 1 above, measurement uncertainty is 0.11.

3. Precision: Using the data from Table 2, there was no significant difference between the plate counts and the values generated with DNA probe (p=0.58). The difference in variance is not significant (p=0.48) for any platform/gene target combination.

4. Recovery: The average of plate counts was 3.40 log the average (adjusted for sample blanks) from DNA probe was 3.65 log. Using this data, the Recovery of the methods was determined to be 107% on both platforms for both gene targets.

Table 2. Data for determination of Precision and Recovery			
Sample	Aliquot	Plate Count (log CFU)	Probe Result (log CFU/g)
1	Blank	N/A	<1.00
1	2X	5.18	4.76
1	2Z	5.18	5.38
1	4X	3.18	2.48
1	4Z	3.18	2.85
1	6X	1.18	<1.00
1	6Z	1.18	1.00
3	Blank	N/A	<1.00
3	2X	5.15	4.65
3	2Z	5.15	4.76
3	4X	3.15	2.90
3	4Z	3.15	3.04
3	6X	1.15	<1.00
3	6Z	1.15	1.78
5	Blank	N/A	1.85
5	2X	5.23	3.54
5	2Z	5.23	3.84
5	4X	3.23	2.70
5	4Z	3.23	3.00
5	6X	1.23	1.48
5	6Z	1.23	1.30
7	Blank	N/A	1.00
7	2X	5.76	5.59
7	2Z	5.76	5.23
7	4X	3.76	3.57
7	4Z	3.76	3.64
7	6X	1.76	1.78
7	6Z	1.76	1.7
9	Blank	N/A	<1.00
9	2X	5.68	5.25
9	2Z	5.68	5.11
9	4X	3.68	3.59
9	4Z	3.68	3.54
9	6X	1.68	2.94
9	6Z	1.68	1.48

5. Specificity: Samples were prepared as above and the interfering organism was spiked at an ~4 log higher concentration than *Vibrio parahaemolyticus*. Using the data from Table 3, the average Specificity of the method is 1.38, which is within the 95% confidence interval of the method (0.44) from 1.

Table 3. Data for determination of Specificity.	
	Probe Result

Sample	(log CFU/g)	
	Spiked with Vp only	Spiked with Vp and Vv
6-Blank	<1.00	---
6-6T	1.60	1.30
6-6U	1.30	<1.00
6-6W	1.48	1.48
6-6X	1.00	1.00
6-6Z	1.48	1.48

6. Working and Linear Range: Based on the data presented in Table 4, there is a significant correlation between the plate counts and CFU values by DNA probe ($p < 0.001$). The correlation coefficient is 0.96, demonstrating the linearity of the method.

Table 4. Data for determination of Working and Linear Range, Limit of Detection, and Limit of Quantitation/Sensitivity

Sample	Aliquot	Plate Count (log CFU)	Probe Result (log CFU/g)
1	1X	6.18	5.36
1	1Z	6.18	6.18
1	2X	5.18	4.76
1	2Z	5.18	5.38
1	4X	3.18	2.48
1	4Z	3.18	2.85
1	6X	1.18	<1.00
1	6Z	1.18	1.00
1	7X	0.18	<1.00
1	7X	0.18	<1.00
3	1X	6.15	6.29
3	1Z	6.15	6.09
3	2X	5.15	4.65
3	2Z	5.15	4.76
3	4X	3.15	2.90
3	4Z	3.15	3.04
3	6X	1.15	<1.00
3	6Z	1.15	1.78
3	7X	0.15	1.00
3	7Z	0.15	<1.00
5	1X	6.23	5.57
5	1Z	6.23	5.64
5	2X	5.23	3.54
5	2Z	5.23	3.84
5	4X	3.23	2.70
5	4Z	3.23	3.00
5	6X	1.23	1.48

5	6Z	1.23	1.30
5	7X	0.23	1.30
5	7Z	0.23	1.48
7	1X	6.76	6.68
7	1Z	6.76	6.37
7	2X	5.76	5.59
7	2Z	5.76	5.23
7	4X	3.76	3.57
7	4Z	3.76	3.64
7	6X	1.76	1.78
7	6Z	1.76	1.70
7	7X	0.76	1.00
7	7Z	0.76	<1.00
9	1X	6.68	6.44
9	1Z	6.68	4.70
9	2X	5.68	5.25
9	2Z	5.68	5.11
9	4X	3.68	3.59
9	4Z	3.68	3.54
9	6X	1.68	2.94
9	6Z	1.68	1.48
9	7X	0.68	<1.00
9	7Z	0.68	<1.00

7. Limit of Detection: The *Limit of Detection of the method is 10 CFU/g*. This is reliant upon the amount of sample (0.1g) that can be tested by the spread plate method.

8. Limit of Quantification/ Sensitivity: The limit of quantification/sensitivity is also reliant upon the amount of sample that can be tested.

9. Ruggednes: Replicate spiked aliquots from each sample were processed with different batches of media/ lots of reagents at the same time. Different samples were processed on different days. Using the data in Table 5, there was *no significant difference (p=0.94) between batches/lots* of media and reagents.

Sample	Probe Result (log CFU/g)	
	Replicate 1 (X)	Replicate 2 (Z)
2	4.78	4.88
4	2.85	2.78

6	1.00	1.48
8	5.64	5.73
10	3.72	3.57

10. Matrix Effects: Effects of oyster matrix on the performance of the method was taken into consideration in testing all of the above criteria by using the sample blank.

11. Additional Data: Inclusivity/Exclusivity. Control filters with the isolates listed below were prepared and tested as outlined above. All isolates. All isolates gave the expected reaction, demonstrating 100% Inclusivity/Exclusivity.

Species	Number of Strains Tested	Number <i>trh</i>-positive
<i>V. parahaemolyticus</i> [†]	43	43
<i>V. parahaemolyticus</i> *	39	0
<i>V. cholerae</i>	25	0
<i>V. vulnificus</i>	13	0
<i>V. metschnikovii</i>	12	0
<i>V. fluvialis</i>	6	0
<i>V. hollisae</i>	5	0
<i>V. algenolyticus</i>	2	0
<i>Salmonella spp.</i>	20	0
<i>Listeria spp.</i>	20	0
Other non- <i>Vibrio</i> species	15	0

[†] *V. parahaemolyticus* strains previously determined to be *trh*-positive.

* *V. parahaemolyticus* strains previously determined to be *trh*-negative.

Step-by-step procedure including equipment, reagents and safety requirements necessary to run the method:

1. Special Equipment, Media, and Reagents
 - 1.1. Special Equipment and Materials Required
 - 1.1.1. Shaking water bath(s) (42°C and 54°C)
 - 1.1.2. Orbital shaker

- 1.1.3. Microwave
- 1.1.4. Plastic tubs with lids (300-500 ml capacity)
- 1.1.5. Whatman 541 filters, 85mm
- 1.1.6. Sterile spread rods
- 1.1.7. Sterile inoculating loops
- 1.1.8. Sterile toothpicks
- 1.1.9. Whirl-Pak bags (4.5"x9")
- 1.2. Media and Reagents
 - 1.2.1. Alkaline peptone water (APW)
 - 1.2.2. Phosphate buffered saline (PBS)
 - 1.2.3. Thiosulfate citrate bile salts sucrose (TCBS) agar
 - 1.2.4. T₁N₃ agar
 - 1.2.5. Lysis solution
 - 1.2.6. 2M ammonium acetate
 - 1.2.7. 20X SSC and 1X SSC
 - 1.2.8. 1X SSC/SDS
 - 1.2.9. Proteinase K
 - 1.2.10. Hybridization solution
 - 1.2.11. NBT/BCIP tablets
 - 1.2.12. AP-labeled DNA probes (DNA Technology)
- 2. Outlined Procedure
 - 2.1. Preparation of shellfish
 - 2.1.1. Hands of examiner must be scrubbed thoroughly with soap and potable water; latex or nitrile gloves should be worn while cleaning oysters.
 - 2.1.2. Scrape off growth and loose material from shell, and scrub shell stock with sterile stiff brush under running water.
 - 2.1.3. Place clean shellstock on clean towels or absorbent paper.
 - 2.1.4. Change gloves and brushes between samples.
 - 2.1.5. Protective chain mail glove can be used under a latex glove; outer gloves should be changed between samples.
 - 2.1.6. Tare a sterile blender.
 - 2.1.7. Using a sterile oyster knife, insert the point between the shells on the ventral side, about ¼ the distance from the hinge to the bill; alternately, knife can be inserted after making small opening with sterile bone cutting forceps.
 - 2.1.8. Cut adductor muscle from upper flat shell and pry the shell wide enough to drain shell liquor into the blender.
 - 2.1.9. The upper shell can then be pried loose at hinge and discarded.
 - 2.1.10. The whole animal (including adductor muscle) should be transferred to the sterile blender after severing the adductor muscle connection to the lower shell.
 - 2.1.11. A minimum of 12 animals or 200g is required.
 - 2.1.12. Blend without adding diluent for 60-120 sec at 14,000 rpm.
 - 2.2. Preparation of spread plates
 - 2.2.1. Prepare 10-fold serial dilutions of shellfish homogenate in PBS
 - 2.2.2. Inoculate 100µl of appropriate dilutions onto pre-dried T₁N₃ agar plates
 - 2.2.3. Spread inoculum gently into agar until completely absorbed
 - 2.2.4. Invert plates and incubate at 30-37°C overnight.

2.2.5. Alternately, this method can be utilized with suspect isolates replicated to T₁N₃ agar from 96 well plates obtained from a standard MPN method

2.3. DNA Probe Colony Hybridization

2.4. Alkaline phosphatase-labeled oligonucleotide probes (*AP-tlh*, *AP-tdh*, and *AP-trh*) can be stored in the refrigerator (4°C) for 1-2 years; do not freeze.

2.5. Filter Preparation

2.5.1. Label #541 Whatman filters with sample number, date, analyst initials, and probe to be hybridized with (*tlh*, *tdh*, or *trh*). Make sure orientation of filter is noted so that positive spots can be correlated to the appropriate well in the microtiter plate. A dot near the A1 well is sufficient.

2.5.2. Place each filter label-side down on appropriate T₁N₃ plate; apply gentle pressure to ensure contact with each colony. Allow labeled filter to sit at RT for 1-30 min. Transfer each filter with colony-side up to a plastic or glass petri dish lid containing 1 ml of lysis solution.

2.5.3. Microwave filters in petri dishes (full power) for 15-20 sec/filter depending on wattage of microwave; rotate dishes with filters and repeat microwaving. Filters should be hot and almost completely dry but not brown.

2.5.4. Transfer filters to a plastic wash container (up to 30 filters can be combined in one container) and neutralize with ammonium acetate (4 ml/filter) for 5 min on shaker at RT.

2.5.5. Decant ammonium acetate and rinse filters 2 times with 1X SSC buffer (10 ml/filter), for 2 min each time. (Filters can be air dried and stored at this point.)

2.6. Proteinase K (proK) treatment

2.6.1. Prepare proK solution (this is made by adding 10 ml/filter of 1X SSC and 20 µl/filter of proK stock solution) for the appropriate number of filters. Place filters (up to 30) in plastic wash container of proK solution. Incubate for 30 min in a 42°C water bath with shaking (50 rpm) to destroy naturally occurring alkaline-phosphatase and digest bacterial protein.

2.6.2. Decant proK solution. Rinse filter 3 times in 1X SSC (10 ml/filter) for 10 min at RT with shaking at 50 rpm. (Filters can be air dried by placing on paper towels and stored when completely dry.)

2.7. Hybridization

2.7.1. Place up to 5 proK-treated filters (either dried or straight from treatment) in a Whirl-Pak bag. Add 10 ml of pre-warmed hybridization buffer and close bag to exclude air. Avoid trapping air bubbles. Incubate filters for 30 min at 54°C in a shaking (50 rpm) water bath.

2.7.2. Pour off buffer from bag and add 10 ml fresh pre-warmed buffer/bag. Add probe (final conc. is 0.5 pmol/ml) to bag with filters. Reseal bag, excluding air, and incubate 1 h in a 54°C water bath with shaking. The temperature is critical for hybridization and washing steps.

2.7.3. Remove filters from hybridization bags and place in plastic wash container(s).

2.7.4. Add 10 ml/filter 1X SSC/1% SDS. Incubate in a 54°C water bath with shaking for 10 min. Repeat wash a second time.

2.7.5. Rinse filter 5 times for 5 min each in 1X SSC at RT on an orbital shaker, 100 rpm.

2.8. Color development

- 2.8.1. In petri dish, add 20 ml of NBT/BCIP solution. Add filters (5 or fewer) to dish and incubate with gentle shaking at 35-37°C; cover to omit light. Check development of positive control every 30 min.
- 2.8.2. Transfer filters to a plastic wash container and add tap water (10 ml/filter). Rinse filters at RT with shaking for 10 min. Repeat rinse 2 additional times to stop color development. Do not expose filters to light as they will continue to develop. Consider purple or brown spots positive.

Validation Data for Direct Plating Method for *trh*, Proposal 15-112

Name of Method Submitter: Jessica L. Jones, Ph.D.

Specific purpose or intent of the method for use in the NSSP: Seeking approval for this method as an approved limited use method that can be used as appropriate for PHP validation and verification testing of oysters, as well as environmental testing such as that which may be required for the re-opening of growing areas closed due to illness.

Validation Criteria Data: For evaluation of all validation criteria below, PHP oysters were obtained in the best effort to find samples free of the target organism. A different lot of PHP oysters was used for each sample. For each sample, a minimum of 10 animals were used to prepare a homogenate. The homogenate was then aliquoted and appropriate aliquots spiked with a *tdh+/trh+ Vibrio parahaemolyticus* culture (unless otherwise noted), while one aliquot was left un inoculated (sample blank). Spike levels were determined by spread plating dilutions of the culture in triplicate onto TSA+2% NaCl. Appropriate aliquots of spiked samples were spread plated onto T1N3 agar and colony lifts hybridized with an alkaline phosphatase-labeled probe specific for *trh* as detailed in the step-by-step procedure. Data were analyzed as described in the “SLV Documents for Marine Biotoxin and Non-MPN Based Microbiological Methods” on the ISSC website.

Table 1. Data for determination of Accuracy/Trueness and Measurement Uncertainty			
Sample	Plate Count (log CFU/g)	Sample Blank Probe Result (log CFU/g)	Spiked Sample Probe Result (log CFU/g)
1-2X	5.18	<1.00	4.76
2-2X	5.18	<1.00	4.78
3-4X	3.15	<1.00	2.90
4-4X	3.15	<1.00	2.85
5-6X	1.15	1.85	1.48
6-6X	1.15	<1.00	1.00
7-2X	5.76	1.00	5.59
8-2X	5.76	<1.00	5.64
9-4X	3.68	<1.00	3.59
10-4X	3.68	<1.00	3.72
11-6X	1.60	<1.00	1.70
12-6X	1.60	<1.00	1.85
13-2X	5.72	<1.00	5.70
14-2X	5.72	<1.00	5.44
15-4X	3.62	<1.00	3.49
16-4X	3.62	<1.00	3.53
17-6X	1.74	1.00	1.70
18-6X	1.62	1.00	2.04
19-2X	5.74	<1.00	5.45
20-6X	1.62	1.00	1.78

1. Accuracy/Trueness: Using the data from Table 1, the average of plate counts was 3.52 log and the average from DNA probe (after adjustment with sample blank results) was 3.48 log. The Accuracy/Trueness of the method is 99%.

Measurement Uncertainty: Using the data from Table 1 above, measurement uncertainty is 0.13 log CFU/g.

Sample	Probe Result (log CFU/g)	
	Replicate 1 (X)	Replicate 2 (Z)
2	4.78	4.88
4	2.85	2.78
6	1.00	1.48
8	5.64	5.73
10	3.72	3.57
12	1.85	1.85
14	5.44	5.29
16	3.53	3.71
18	2.04	2.08
20	1.78	1.78

2. Ruggedness: Replicate spiked aliquots from each sample were processed with different batches of media/ lots of reagents at the same time. Different samples were processed on different days. Using the data in Table 2, a left skew of both sets of data was observed, with a variance ratio of 1.09 (not significant, $p=0.91$), so a paired t-test was used to compare the results. There was no significant difference ($p=0.94$) between batches/lots of media and reagents.

Sample	Aliquot	Plate Count (log CFU/g)	Sample Blank Probe Result (log CFU/g)	Spiked Sample Probe Result (log CFU/g)
1	2x	5.18	<1.00	4.76
1	2z	5.18	<1.00	5.38
1	4x	3.18	<1.00	2.48
1	4z	3.18	<1.00	2.85
1	6x	1.18	<1.00	<1.00
1	6z	1.18	<1.00	1.00
3	2x	5.15	<1.00	4.65
3	2z	5.15	<1.00	4.76
3	4x	3.15	<1.00	2.90
3	4z	3.15	<1.00	3.04
3	6x	1.15	<1.00	<1.00
3	6z	1.15	<1.00	1.78
5	2x	5.16	1.85	3.54
5	2z	5.16	1.85	3.84
5	4x	3.16	1.85	2.70
5	4z	3.16	1.85	3.00

5	6x	1.15	1.85	1.48
5	6z	1.15	1.85	1.30
7	2x	5.76	1.00	5.59
7	2z	5.76	1.00	5.23
7	4x	3.76	1.00	3.57
7	4z	3.76	1.00	3.64
7	5x	2.76	1.00	2.58
7	5z	2.76	1.00	2.57
7	6x	1.76	1.00	1.78
7	6z	1.76	1.00	1.70
9	2x	5.68	<1.00	5.25
9	2z	5.68	<1.00	5.11
9	4x	3.68	<1.00	3.59
9	4z	3.68	<1.00	3.54
9	6x	1.68	<1.00	2.94
9	6z	1.68	<1.00	1.48
11	2x	5.60	<1.00	5.54
11	2z	5.60	<1.00	5.46
11	4x	3.60	<1.00	3.48
11	4z	3.60	<1.00	3.71
11	6x	1.60	<1.00	1.70
11	6z	1.60	<1.00	1.00
13	2x	5.72	<1.00	5.70
13	2z	5.72	<1.00	5.08
13	4x	3.72	<1.00	3.36
13	4z	3.72	<1.00	3.36
13	6x	1.72	<1.00	1.48
13	6z	1.72	<1.00	1.30
15	2x	5.62	<1.00	5.53
15	2z	5.62	<1.00	4.98
15	4x	3.62	<1.00	3.49
15	4z	3.62	<1.00	3.48
15	6x	1.62	<1.00	1.30
15	6z	1.62	<1.00	1.00
17	2x	5.74	1.00	5.51
17	2z	5.74	1.00	5.42
17	4x	3.74	1.00	3.57
17	4z	3.74	1.00	3.64
17	6x	1.74	1.00	1.70
17	6z	1.74	1.00	1.90
19	2x	5.74	<1.00	5.45

19	2z	5.74	<1.00	5.64
19	4x	3.74	<1.00	3.51
19	4z	3.74	<1.00	3.45
19	6x	1.74	<1.00	1.78
19	6z	1.74	<1.00	1.70

3. Precision: Using the data from Table 3, there was no significant difference ($p>0.05$) in the variance ratio across the range of concentrations, with a coefficient of variance of 45% for the method. Additionally, there was no significant difference between the plate counts and the values generated with DNA probe ($p=0.68$).

Recovery: The average of plate counts was 3.51 log CFU/g the average (adjusted for sample blanks) from DNA probe was 3.28 log CFU/g. Using this data, the Recovery of the method was determined to be 93%.

Table 4. Data for determination of Specificity				
Sample	Aliquot (Vp only)	Log CFU/g	Aliquot (Vp and Vv)	Log CFU/g
6	6T	1.60	VVT	1.30
6	6U	1.30	VVU	<1.00
6	6W	1.48	VVW	1.48
6	6X	1.00	VVX	1.00
6	6Z	1.48	VVZ	1.48
Sample	Aliquot (Vp only)	Log CFU/g	Aliquot (Vp and Vc)	Log CFU/g
12	6T	1.60	VCT	1.48
12	6U	2.00	VCU	<1.00
12	6W	1.70	VCW	1.60
12	6X	1.85	VCX	1.48
12	6Z	1.85	VCZ	1.00
Sample	Aliquot (Vp only)	Log CFU/g	Aliquot (Vp and Vf)	Log CFU/g
18	6T	1.95	VFT	<10.00
18	6U	2.11	VFU	<10.00
18	6W	2.11	VFW	<10.00
18	6X	2.04	VFX	<10.00
18	6Z	2.08	VFZ	<10.00
Sample	Aliquot (Vp only)	Log CFU/g	Aliquot (Vp and Va)	Log CFU/g
20	6T	1.70	VAT	<1.00
20	6U	1.48	VAU	<1.00
20	6W	1.78	VAW	<1.00
20	6X	1.78	VAX	<1.00
20	6Z	1.78	VAZ	<1.00

4. Specificity: Interfering organisms tested were *V. vulnificus* (Vv), *V. cholerae* (Vc), *V. fluvialis* (Vf), and *V. alginolyticus* (Va). Using the data from Table 4, the overall average Specificity Index of the method is 1.43, which is within the 95% confidence interval of the method (0.44) from 1.

However, significant differences between the average specificity indices and 1 were observed when examining the data from each interfering organism. Differences were significant for *V. fluvialis* ($p < 0.001$) and *V. alginolyticus* ($p < 0.001$), as well as *V. cholerae* ($p = 0.05$). This significance is likely due to the differences in spike levels, rather than the specific organism as the ratio of *V. parahaemolyticus* to interfering organism was 1:13000, 1:6000, 1:1500, and 1:240 for *V. fluvialis*, *V. alginolyticus*, *V. cholerae*, and *V. vulnificus*, respectively. Additionally, it should be noted that filters were lifted from plates with more colonies than recommended from the samples with the interfering organisms to give the best likelihood of enumerating *V. parahaemolyticus*. Together, this indicates that the method specificity is dependent on the ratio of target to interfering organism, where ratios of 1:1000 may cause interference, but lower ratios do not. In real world samples, this ratio of *trh+* *V. parahaemolyticus* to other *Vibrio* species, is unlikely, supporting the fit-for-purpose of this method.

Table 5. Data for Working/Linear Range			
Sample	Plate Count	Replicate 1 (X)	Replicate 2 (Z)
1-1	6.18	5.36	6.18
1-2	5.18	4.76	5.38
1-4	3.18	2.48	2.85
1-5	2.18	2.00	1.85
1-6	1.18	<1.00	1.00
1-7	0.18	<1.00	<1.00
3-1	6.15	6.29	6.09
3-2	5.15	4.65	4.76
3-4	3.15	2.90	3.04
3-5	2.15	2.23	2.68
3-6	1.15	<1.00	1.78
3-7	0.15	1.00	<1.00
5-1	6.16	5.57	5.64
5-2	5.16	3.54	3.84
5-4	3.16	2.70	3.00
5-5	2.16	2.20	2.38
5-6	1.15	1.48	1.30
5-7	0.15	1.30	1.48
7-1	6.76	6.68	6.37
7-2	5.76	5.59	5.23
7-4	3.76	3.57	3.64
7-5	2.76	2.58	2.57
7-6	1.76	1.78	1.70

7-7	0.76	1.00	<1.00
9-1	6.68	6.44	4.70
9-2	5.68	5.25	5.11
9-4	3.68	3.59	3.54
9-5	2.68	2.49	2.79
9-6	1.68	2.94	1.48
9-7	0.68	<1.00	<1.00
11-1	6.60	6.40	6.44
11-2	5.60	5.54	5.46
11-4	3.60	3.48	3.71
11-5	2.60	3.06	2.84
11-6	1.60	1.70	1.00
11-7	0.60	<1.00	<1.00
13-1	6.72	6.71	5.55
13-2	5.72	5.70	5.08
13-4	3.72	3.36	3.36
13-5	2.72	2.48	2.38
13-6	1.72	1.48	1.30
13-7	0.72	<1.00	<1.00
15-1	6.62	6.22	6.40
15-2	5.62	5.53	4.98
15-4	3.62	3.49	3.48
15-5	2.62	1.85	1.60
15-6	1.62	1.30	1.00
15-7	0.62	<1.00	<1.00
17-1	6.74	6.33	6.39
17-2	5.74	5.51	5.42
17-4	3.74	3.57	3.64
17-5	2.74	2.60	2.72
17-6	1.74	1.70	1.90
17-7	0.74	1.30	<1.00
19-1	6.74	6.36	6.32
19-2	5.74	5.45	5.64
19-4	3.74	3.51	3.45
19-5	2.74	2.72	2.53
19-6	1.74	1.78	1.70
19-7	0.74	1.30	<1.00

5. Working and Linear Range: Based on the data presented in Table 4, the linear range of the method is 50 to 100,000 CFU/g. There is a significant correlation between the plate counts and CFU values by DNA probe ($p < 0.001$), with a correlation coefficient is 0.93.

Limit of Detection: The theoretical LOD based on the data above is 4.7 CFU/g. However, this is reliant upon the amount of sample (0.1g) that can be tested by the spread plate method. Therefore, the Limit of Detection of the method is 10 CFU/g.

Limit of Quantitation/ Sensitivity: The limit of quantification/sensitivity is also reliant upon the amount of sample that can be tested. As such, the Limit of Quantitation of the method is 10 CFU/g.

Additional Data: Inclusivity/Exclusivity. Control filters with the isolates listed below were prepared and tested as outlined above. All isolates. All isolates gave the expected reaction, demonstrating 100% Inclusivity/Exclusivity.

Species	Number of Strains Tested	Number <i>trh</i> -positive
<i>V. parahaemolyticus</i> [†]	43	43
<i>V. parahaemolyticus</i> *	39	0
<i>V. cholerae</i>	25	0
<i>V. vulnificus</i>	13	0
<i>V. metschnikovii</i>	12	0
<i>V. fluvialis</i>	6	0
<i>Grimontia hollisae</i>	5	0
<i>V. alginolyticus</i>	2	0
<i>Salmonella spp.</i>	20	0
<i>Listeria spp.</i>	20	0
Other non- <i>Vibrio</i> species	15	0

[†] *V. parahaemolyticus* strains previously determined to be *trh*-positive.

* *V. parahaemolyticus* strains previously determined to be *trh*-negative.

Step-by-step procedure including equipment, reagents and safety requirements necessary to run the method:

1. Special Equipment, Media, and Reagents
 - 1.1. Special Equipment and Materials Required
 - 1.1.1. Shaking water bath(s) (42°C and 54°C)

- 1.1.2.Orbital shaker
- 1.1.3.Microwave
- 1.1.4.Plastic tubs with lids (300-500 ml capacity)
- 1.1.5.Whatman 541 filters, 85mm
- 1.1.6.Sterile spread rods
- 1.1.7.Sterile inoculating loops
- 1.1.8.Sterile toothpicks
- 1.1.9.Whirl-Pak bags (4.5"x9")
- 1.2. Media and Reagents
 - 1.2.1.Alkaline peptone water (APW)
 - 1.2.2.Phosphate buffered saline (PBS)
 - 1.2.3.Thiosulfate citrate bile salts sucrose (TCBS) agar
 - 1.2.4.T₁N₃ agar
 - 1.2.5.Lysis solution
 - 1.2.6.2M ammonium acetate
 - 1.2.7.20X SSC and 1X SSC
 - 1.2.8.1X SSC/SDS
 - 1.2.9.Proteinase K
 - 1.2.10. Hybridization solution
 - 1.2.11. NBT/BCIP tablets
 - 1.2.12. AP-labeled DNA probes (DNA Technology)
- 2. Outlined Procedure
 - 2.1. Preparation of shellfish
 - 2.1.1. Hands of examiner must be scrubbed thoroughly with soap and potable water; latex or nitrile gloves should be worn while cleaning oysters.
 - 2.1.2. Scrape off growth and loose material from shell, and scrub shell stock with sterile stiff brush under running water.
 - 2.1.3. Place clean shellstock on clean towels or absorbent paper.
 - 2.1.4. Change gloves and brushes between samples.
 - 2.1.5. Protective chain mail glove can be used under a latex glove; outer gloves should be changed between samples.
 - 2.1.6. Tare a sterile blender.
 - 2.1.7. Using a sterile oyster knife, insert the point between the shells on the ventral side, about ¼ the distance from the hinge to the bill; alternately, knife can be inserted after making small opening with sterile bone cutting forceps.
 - 2.1.8. Cut adductor muscle from upper flat shell and pry the shell wide enough to drain shell liquor into the blender.
 - 2.1.9. The upper shell can then be pried loose at hinge and discarded.
 - 2.1.10.The whole animal (including adductor muscle) should be transferred to the sterile blender after severing the adductor muscle connection to the lower shell.
 - 2.1.11.A minimum of 12 animals or 200g is required.
 - 2.1.12.Blend without adding diluent or with equal weight of diluent (APW or PBS) for 60-120 sec at 14,000 rpm.
 - 2.2. Preparation of spread plates
 - 2.2.1.Prepare 10-fold serial dilutions of shellfish homogenate in PBS
 - 2.2.2.Inoculate 100µl of appropriate dilutions onto pre-dried T₁N₃ agar plates
 - 2.2.3.Spread inoculum gently into agar until completely absorbed

- 2.2.4. Invert plates and incubate at 30-37°C overnight.
- 2.2.5. Alternately, this method can be utilized with suspect isolates replicated to T₁N₃ agar from 96 well plates obtained from a standard MPN method
- 2.3. DNA Probe Colony Hybridization
 - 2.4. Alkaline phosphatase-labeled oligonucleotide probes (*AP-tlh*, *AP-tdh*, and *AP-trh*) can be stored in the refrigerator (4°C) for 1-2 years; do not freeze.
 - 2.5. Filter Preparation
 - 2.5.1. Label #541 Whatman filters with sample number, date, analyst initials, and probe to be hybridized with (*tlh*, *tdh*, or *trh*). Make sure orientation of filter is noted so that positive spots can be correlated to the appropriate well in the microtiter plate. A dot near the A1 well is sufficient.
 - 2.5.2. Place each filter label-side down on appropriate T₁N₃ plate; apply gentle pressure to ensure contact with each colony. Allow labeled filter to sit at RT for 1-30 min. Transfer each filter with colony-side up to a plastic or glass petri dish lid containing 1 ml of lysis solution. Alternatively, to avoid overheating, a Whatman #3 filter pad can be saturated with 4ml of lysis buffer in the bottom of the petri dish onto which the #541 filter is transferred.
 - 2.5.3. Microwave filters in petri dishes (full power) for 15-20 sec/filter depending on wattage of microwave; rotate dishes with filters and repeat microwaving. Filters should be hot and almost completely dry but not brown.
 - 2.5.4. Transfer filters to a plastic wash container (up to 30 filters can be combined in one container) and neutralize with ammonium acetate (4 ml/filter) for 5 min on shaker at RT.
 - 2.5.5. Decant ammonium acetate and rinse filters 2 times with 1X SSC buffer (10 ml/filter), for 1-2 min each time. (Filters can be air dried and stored at this point.)
 - 2.6. Proteinase K (proK) treatment
 - 2.6.1. Prepare proK solution (this is made by adding 10 ml/filter of 1X SSC and 20 µl/filter of proK stock solution) for the appropriate number of filters. Place filters (up to 30) in plastic wash container of proK solution. Incubate for 30 min in a 42°C water bath with shaking (50 rpm) to destroy naturally occurring alkaline-phosphatase and digest bacterial protein.
 - 2.6.2. Decant proK solution. Rinse filter 3 times in 1X SSC (10 ml/filter) for 10 min at RT with shaking at 50 rpm. (Filters can be air dried by placing on paper towels and stored when completely dry.)
- 2.7. Hybridization
 - 2.7.1. Place up to 5 proK-treated filters (either dried or straight from treatment) in a Whirl-Pak bag. Add 10 ml of pre-warmed hybridization buffer and close bag to exclude air. Avoid trapping air bubbles. Incubate filters for 30 min at 54°C in a shaking (50 rpm) water bath.
 - 2.7.2. Pour off buffer from bag and add 10 ml fresh pre-warmed buffer/bag. Add probe (final conc. is 0.5 pmol/ml) to bag with filters. Reseal bag, excluding air, and incubate 1 h in a 54°C water bath with shaking. The temperature is critical for hybridization and washing steps.
 - 2.7.3. Remove filters from hybridization bags and place in plastic wash container(s).

- 2.7.4. Add 10 ml/filter 1X SSC/1% SDS . Incubate in a 54°C water bath with shaking for 10 min. Repeat wash a second time.
- 2.7.5. Rinse filter 5 times for 5 min each in 1X SSC (10ml/ filter) at RT on an orbital shaker, 100 rpm.
- 2.8. Color development
 - 2.8.1. In petri dish, add 20 ml of NBT/BCIP solution. Add filters (5 or fewer) to dish and incubate with gentle shaking; cover to omit light. Incubation temperatures from room temperature up to 40°C can be used; color development will be quicker at higher temperatures. Check development of positive control every hour.
 - 2.8.2. Transfer filters to a plastic wash container and add tap water (10 ml/filter). Rinse filters at RT with shaking for 10 min. Repeat rinse 2 additional times to stop color development. Do not expose filters to light as they will continue to develop. Consider purple or brown spots positive.

Method Application and Single Lab Validation Checklist For Acceptance of a Method for Use in the NSSP

The purpose of single laboratory validation in the National Shellfish Sanitation Program (NSSP) is to ensure that the analytical method under consideration for adoption by the NSSP is fit for its intended use in the Program. A Checklist has been developed which explores and articulates the need for the method in the NSSP; provides an itemized list of method documentation requirements; and, sets forth the performance characteristics to be tested as part of the overall process of single laboratory validation. For ease in application, the performance characteristics listed under validation criteria on the Checklist have been defined and accompany the Checklist as part of the process of single laboratory validation. Further a generic protocol has been developed that provides the basic framework for integrating the requirements for the single laboratory validation of all analytical methods intended for adoption by the NSSP. Methods submitted to the ISSC LMR Committee for acceptance will require at a minimum 6 months for review from the date of submission.

Name of the New Method	Male-specific Coliphage for Wastewater
Name of the Method Developer	USFDA Gulf Coast Seafood Laboratory
Developer Contact Information	USFDA Gulf Coast Seafood Laboratory, 1 Iberville Drive, Dauphin Island, AL 36 kevin.calci@fda.hhs.gov

Checklist	Y/N	Submitter Comments
A. Need for the New Method		
Clearly define the need for which the method has been developed.	Y	
What is the intended purpose of the method?	Y	
Is there an acknowledged need for this method in the NSSP?	Y	
What type of method? i.e. chemical, molecular, culture, etc.	Y	Culture method for Male-specific Coliphage in Wastewater Influent, Mid-process Samples, and Effluent

B. Method Documentation		
1. Method documentation includes the following information:		
Method Title	Y	
Method Scope	Y	
References	Y	
Principle	Y	
Any proprietary aspects	N	
Equipment required	Y	
Reagents required	Y	
Sample collection, preservation and storage requirements	Y	

Safety requirements	Y	
Clear and easy to follow step-by-step procedure	Y	
Quality control steps specific for this method	Y	

C. Validation Criteria		
1. Accuracy / Trueness	Y	
2. Measurement uncertainty	Y	
3. Precision characteristics (repeatability)	Y	
4. Recovery	Y	
5. Specificity	NA	
6. Working and Linear ranges	Y	
7. Limit of detection	Y	
8. Limit of quantitation / Sensitivity	Y	
9. Ruggedness	Y	
10. Matrix effects	N	
11. Comparability (if intended as a substitute for an established method accepted by the NSSP)	NA	

D. Other Information		
1. Cost of the method	Y	
2. Special technical skills required to perform the method	Y	
3. Special equipment required and associated cost	Y	
4. Abbreviations and acronyms defined	Y	
5. Details of turn around times (time involved to complete the method)	Y	
6. Provide brief overview of the quality systems used in the lab	Y	

Submitters Signature	Date:
Submission of validation data and draft method to committee	Date:
Reviewing members:	
Accepted	Date:
Recommendations for further work	Date:

Comments:

**Single Laboratory Validation (SLV) Protocol For Submission
to the ISSC For Method Approval**

Name of the New Method - A Culture Method/Double Agar Overlay Method for the Determination of Male-specific Coliphage (MSC) for Wastewater

Name of Method Developer - Kevin Calci, USFDA-GCSL

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Date of Interim Submission - May 15, 2017

Section A. Need for the New Method

FDA has long been using Male-Specific Coliphage (MSC) to evaluate the potential viral contamination of shellfish growing water by wastewater treatment plant (WTP) outfalls. Methods using MSC as an indicator of viral contamination have been successful in evaluation of viral persistence in molluscan shellfish impacted by WTP outfalls (Daskin et al, 2008)(ISSC MSC Workshop). Studies continue to show a significant inverse relationship between decreasing MSC levels in shellfish and increasing wastewater dilution, which is in turn strongly associated with increasing distance from the WTP discharges (Goblic et al, 2011). The relationship between the level of viral contamination in shellfish and dilution of treated wastewater is really contingent on the viral reduction efficiency of the WTP impacting the area.

The purpose of this method is to assess the log₁₀ reduction of MSC, as a process indicator for enteric viruses, namely Human Norovirus, in wastewater samples including raw influent, pre-disinfected effluent and final effluent. By comparing log₁₀ values of these results, the viral reduction performance of a WTP can be assessed under different environmental and operational conditions (Amarasiri et al, 2017) (Pouillot et al, 2015). Understanding the viral reduction performance at different stages in a wastewater treatment process is a valuable assessment tool to determine growing area classification and management options for shellfish growing areas adjacent to and downstream from the WTP outfall. This newly configured FDA method for the determination of MSC in wastewater samples has been adapted from previous methods so that it may be more readily implemented at NSSP Laboratories.

The recognized need for an alternative viral indicator is addressed in detail in the newly accepted 2015 Revision of the NSSP Guide for the Control of Molluscan Shellfish, Section IV Guidance Documents, Chapter II, @ .19, Determining Appropriately Sized Prohibited Areas Associated with Wastewater Treatment Plants, page 292. The need and utility for this method was likewise address at the MSC Informational Meeting of the Growing Area Committee (MSC Summit) in

Charlotte, NC in August 2014. A pre-proposal was reviewed at the 2015 ISSC Meeting and given the Proposal Number of 15-114. The LMRC agreed that the pre-proposal was sufficient and that there is a need for the method. The LMRC recommended to Task Force I that Proposal 15-114 be referred to an appropriate committee as determined by the Conference Chair and await the SLV data.

Section B. Method Documentation

Modified Double Agar Overlay Method for Determination of Male-specific Coliphage in Wastewater

May 2017 Revision

This method for determining levels of male-specific coliphage in wastewater is based on the method described by Cabelli in work on the Narragansett Bay Project. (Cabelli, 1998) The development of an *Escherichia coli* host cell that constitutively expresses the F plasmid and is resistant to 95% of the somatic phage in wastewater was described in a subsequent paper. (DeBartolomeis and Cabelli, 1991) FDA refined the method for oyster and hard clam meats as described in the workshop instructions, *Male-specific Bacteriophage (MSB) Workshop*, conducted in Gloucester, Massachusetts on March 9-12, 2004. (US Food and Drug Administration, 2004) This original FDA (2004) method was submitted as ISSC Proposal 05-114. Spinney Creek Shellfish, Inc. (SCS) further refined these procedures for soft-shelled clams and American Oysters in work funded by the Maine Technology Institute in 2006 with the assistance of Mercuria Cumbo of the Maine Department of Marine Resources. This method was approved for limited use by the 2009 ISSC in Manchester, NH. In work funded by UNH Sea Grant, SLV work continued for species extension to quahogs, which was approved for limited use by the 2013 ISSC in San Antonio, Texas. Method development and preliminary SLV trials were conducted in 2015 by Kevin Calci and Ashley Cooper at USFDA-GCSL. Additional SLV trials were conducted at the Spinney Creek Shellfish Laboratory in collaboration with Kevin Calci, the method developer in work supported by UNH Sea Grant.

A. Apparatus and Materials.

Equipment and Materials for Collection and Transport of Wastewater Samples:

250 or 500 ml Sterile Sample Containers
 Sealable Bio-hazard Bags (used when shipping)
 Labels
 Cooler
 Gel Packs
 Sampling Device
 10% Sodium Thiosulfate Solution (for effluent samples)

Laboratory Equipment:

Water bath, 50-52°C
 Air Incubator, 35-37°C
 Balance
 Stir plate and magnetic stirring bars, sterile
 Mini vortexer
 Autoclave, 119°C - 121°C
 Refrigerator, 0-4° C
 Freezer, -20°C

pH meter
Erlenmeyer flasks, 2L and 4L
Graduated cylinders, 1000 ml
500 ml jars, autoclavable with caps
Inoculating loops (3 mm in diameter or 10 μ L volume)
Bacti-cinerator or flame
Sterile swabs
Sterile, disposable filters, 0.22 or 0.45 μ m pore size
Syringes, sterile disposable; 5ml
Serological Pipets- 1 ml, 2 ml, 5 ml, 10 ml
Pipet-aid, or
(Micropipette option; 100 μ L and 1000 μ L (marked with red tape for positive controls), 200 μ L (for aliquots of host cells), 2500 μ L (for sample aliquots), Micropipette tips, sterile 100 μ L, 1000 μ L, 2500 μ L Micropipette Stand)
Petri dishes, sterile disposable 100 x 15 mm
Petri dish racks
Test tubes 16 x 100 mm (for soft agar)
Dilution tubes, 16 x 150 mm, sterile with screw caps
50ml conical tubes, sterile with screw caps
Test tube racks--sizes to accommodate tubes
Freezer vials, sterile 30 ml with screw caps
Baskets with tops to hold freezer vials
Parafilm tape
Aluminum foil
Counter-pen, digital

Reagents:

Reagent water
Glycerol- sterile
Ethanol, 70% or laboratory disinfectant
Calcium chloride, 1M
Mineral oil
Sodium Thiosulfate (for effluent sample bottles to eliminate chlorine residual)

Antibiotic stocks:

Ampicillin sodium salt (Sigma A9518)
Streptomycin sulfate (Sigma S6501)
Streptomycin and Ampicillin stock solutions (50 μ g/ml each).
Note: Antibiotics must always be added to liquids and media after these have been autoclaved and cooled.

Media Components:

Agar, Granulated
Dextrose
NaCL

CaCl₂
DI water

Media:

Bottom Agar
DS Soft Agar
Growth Broth

Bacterial Host Strain:

E. coli F_{amp} □ *E. coli* HS(pF_{amp})RR (ATCC # 700891).

MSC (Coliphage) Stock:

Type Strain - MS2, ATCC # 15597

B. Media Composition.

Bottom Agar:

Tryptone	10.0 g
Dextrose	1.0 g
NaCl	5.0 g
Agar	15.0 g
DI water	990 ml
Final pH	6.7 ± 0.2 at 25°C

1. With gentle mixing, add all the components, except antibiotics, to 990ml of dH₂O in a 1000ml flask (increase flask size to make larger volumes). Dissolve, heat until clear, bringing to a boil.
2. Sterilize at 121°C ± 2°C for 15 minutes.
3. Temper to 50°C in the water bath.
4. Add 5 ml of Streptomycin sulfate/Ampicillin solution, aseptically to the flask (50 µg/ml each in final) and mix. Transfer to 2 – 500ml sterile jars (easier to pour plates from jars).
5. Pour 15-17 ml aliquots aseptically into sterile 100 x 15 mm Petri dishes and allow the agar to harden. Tip Petri dish lids off slightly to reduce condensation.
6. Store bottom agar plates inverted at 4°C and warm to room temperature for 1 hour before use.
7. Plates stored sealed at 4°C can be used up to 3 months.

Streptomycin sulfate/Ampicillin Solution:

1. Dissolve 0.5g of streptomycin sulfate and 0.5g of ampicillin in 50 ml of dH₂O with a sterile 100 ml graduated cylinder in sterile 600 ml beaker with sterile stir bar.
2. Stir for 2 to 3 minutes, no heat.
3. Filter by injecting through a sterile 0.22 µm filter.
4. Store in 5 ml aliquots in sterile 30 ml capped freezer vials at -20°C for up to one year. Label and date.
5. Allow to come to room temperature before adding and mixing in tempered bottom agar at 50°C.

DS Soft Agar:

Tryptone	10.0 g
Dextrose	1.0 g
NaCl	5.0 g
1M CaCl ₂	0.5 ml
Agar	7.0 g
DI water	500 ml
Final pH	6.7 ± 0.2

1. With gentle mixing, add all the components to 500 ml of dH₂O in a 1000 ml flask.
2. Bring flask contents to a boil.
3. Dispense in 2.5 ml aliquots into 16 x 100 ml tubes, cover and freeze (-20°C)
4. Sterilize prior to use at 121°C ± 2°C for 15 minutes, then temper to 50-52°C in a water bath set to 50 °C ± 2 °C for no longer than 2 hours.
5. Store up to 3 months at -20 °C.

1M CaCl₂ Solution:

1. Add 11.1 g of CaCl₂ anhydrous (FW 111.0, Dihydrate FW 147) to 100 ml dH₂O in a screw top bottle and dissolve or use prepared from VWR.
2. Sterilize by autoclaving at 121°C for 15 minutes.
3. Store up to three months at 4°C.

Growth Broth:

Tryptone	10.0 g
Dextrose	1.0 g
NaCl	5.0 g
DI water	1000 ml

1. With gentle mixing, add all the components to 1000 ml of dH₂O water in a 2000 ml flask.
2. Dissolve and dispense into sterile screw top containers.
3. Sterilize at 121°C ± 2°C for 15 minutes.
4. Store for up to three months at 4°C.

C. Storage and Propagation of Host Strain, E. coli F_{amp}:**Storage:**

1. Lab stock culture – Frozen at – 80°C indefinitely (most desirable method) in broth culture containing 10% glycerol under no selective pressure. Selective pressure is reapplied when the culture is retrieved, by streaking onto Bottom Agar plates containing the two antibiotics.
2. Long-term working stock culture – Grown tryptic soy agar slant with sterile mineral oil overlay under no selective pressure and stored at room temperature in the dark for up to 2 years.
3. Long-term working stock – 6-hour grown tryptic soy agar slant and deep stab with sterile mineral oil overlay containing the two antibiotics, Ampicillin and Streptomycin (least desirable method).
4. Short-term working stock culture - Grown Bottom Agar streak plate stored at 4°C up to 3 weeks.

Glycerol Solution, 10%:

1. Add 9 ml of distilled water to 1 ml of undiluted glycerol.
2. Autoclave resulting 10% glycerol solution at 121°C for 15 minutes and use at room temperature.
3. For storage, add 1/5th volume of 10% glycerol solution, let stand for 30 minutes, dispense 1 ml aliquots in 2 ml cryo-vials and store at -70 to –80°C (best) or at –20°C.

D. Control Plates.

1. Negative Control - Add 2.5 ml of Growth Broth and 0.2 ml host to the 2.5 ml DS Soft Agar tube.
2. Positive Control - Make serial dilutions using growth broth of the concentrated MS2 control (to grow approximately 50-100 PFU per 2.5 ml), and add 2.5 ml of appropriate MS2 dilution and 0.2 ml of host to 2.5 ml DS Soft agar.

E. MSC Density Determinations in wastewater Treatment Plant (WWTP) Samples.

Sample Requirements:

1. Sterile 250 or 500 ml Nalgene bottles (or comparable bottle) with a permanent fill mark at the approximate 200 or 400 ml level are recommended for wastewater samples including influent, pre-treated effluent, and effluent.
2. Sample collection bottles must be properly labeled with sample number, location, sample type, date and time.
3. Sample bottles are filled to the 200 or 400 ml line. Effluent sample bottles must contain 1.0 ml of 10% sodium thiosulfate solution for 200 ml or 2.0 ml of 10% sodium thiosulfate solution to for 400 ml to inactivate any residual chlorine.

4. Wastewater samples are held under refrigerated conditions at 1-4°C.

Note - A sealed bio-hazard bag is recommended for the samples bottles containing sewage samples going into an insulated shipping box when using overnight carriers.

Propagation of Host Cells:

1. Allow grown Bottom Agar streak plate and Growth Broth to temper to room temperature.
2. Vortex to aerate 20 ml of Growth Broth in a 16 x 150 mm tube, with screw cap.
3. Transfer host strain to Growth Broth using sterile swab to collect material from three colonies off grown Bottom Agar streak plate.
4. Gently shake to mix, then incubate at 35–37°C for 4-6 hours.
5. Once turbidity is observed, use of the host strain broth culture (log-phased growth) may commence.

Note - Following initial inoculation and mixing, do not shake or mix the host strain broth culture (to avoid mixing of cell debris at bottom with log-phase E. coli with pili)

Preparation of Wastewater samples for Analysis:

1. Analyst must wear gloves during handling of stir bars and sample bottle.
2. Water samples are removed from 1 - 4° C.
3. Sample bottle is shaken vigorously for 20 seconds (ensure cap is tightened), and a sterile magnetic stir bar is aseptically transferred to bottle.
4. Sample bottle is placed on stir plate set to medium for five minutes prior to analysis.
5. For the **high range** of this method a 10⁻² decimal dilution is prepared by transferring 1ml of sample with a sterile 2 ml pipette (using a pipette aid) to a sterile 16x150mm screw cap tube containing 9 ml of growth broth. Sample tube is then vortexed for 10 seconds. For the second decimal dilution, 2ml are transferred from the first tube to a sterile 50ml conical tube with cap containing 18ml of growth broth using a second sterile 2ml pipette. The appropriately labeled 50 ml conical tube is then vortexed for 10 seconds.
6. For the **low range** of this method, 30ml of sample is transferred to a sterile 50 mm conical tube with cap using a sterile 10 ml pipette. The appropriately labeled 50 ml conical tube is vortexed for 10 seconds.
7. Prepped samples in labeled 50ml conical tube are stored in a test tube rack which can be stored short term at 0-4°C.
8. Return sample bottles to refrigeration and clean the work surface with disinfectant.

Note: The samples bottles containing wastewater samples should be autoclaved prior to disposal. Sample bottles must be washed and sterilized for re-use.

Direct Analytical Technique for WWTP samples:

This MSC method for wastewater has both a **high range** and a **low range** routine. Combined, the working range is from 5 to 1,200,000 PFU/100ml. The **high range** routine is adequate for enumeration of MSC in WWTP influent and has a working range from 1,000 to 1,200,000 PFU/100ml. The **low range** routine is generally adequate for enumeration of MSC in final effluent and has a working range from 5 to 12,000 PFU/100ml. When testing for pre-treatment effluent (before disinfection) or at times when the effluent is questionable, both high and low ranges routines should be used together.

1. In the morning, propagate host cells as described above.
2. Tubes may be inoculated on a staggered time schedule:

Tubes in incubator at 7:00am	Ready at 11:00am
Tubes in incubator at 8:00am	Ready at 12:00pm
Tubes in incubator at 10:00am	Ready at 2:00pm
Tubes in incubator at 11:00am	Ready at 3:00pm
3. Before experimentation, prepare the wastewater samples for analysis as described above.
4. One hour before experimentation (at 3 hours of host growth), autoclave required number of soft agar tubes at 121°C for 15min. Temper soft agar tubes in water bath set to 50-52°C.

High Range Routine:

For each high range (influent) sample, four (4) Bottom Agar plates and four (4) 2.5 ml DS Soft Agar tubes are prepared. Always begin the day's analyses with a negative control (blank) plate and finish analyses with a positive control plate and a negative control plate.

1. Allow prepared samples (50ml conical tubes, racked and labeled) to warm to room temperature immediately before analysis (20-30 minutes)
2. Remove tubes containing appropriate timed host cells from incubator immediately before analysis.
3. Vortex sample in 50ml conical tube for 10 seconds.
4. Moving quickly and smoothly, gently pipette 200µL of host cells into each of 4 soft agar tubes using a 1 ml serological pipet or 200µL micropipette with sterile tip.
5. Immediately thereafter, pipette 2500µL aliquot of sample into each of the 4 soft tubes using a 10 ml serological pipet and pipet aid or 2500µL micropipette with sterile tip.
6. Mix sample, host and soft agar by gently rolling test tube between hands.

Note: *Once E. coli F_{amp} is added to the mixture do not shake, only gently mix contents by rolling the tube between palms.*

7. Then overlay the contents of the soft agar tubes onto a Bottom Agar plates. Drag the mixture into a clear area and gently swirl the plates to spread sample and agar

mixture.

8. Allow plates to set then inverted and incubated for 16 - 20 hours at 35- 37°C.
9. Circular zones of clearing (typically 1 to 10 mm in diameter) in lawn of host bacteria after 16 – 20 hours of incubation are counted as plaques. PFUs on each plate are counted and totaled using a digital counter pen and adequate light
10. Calculations of **High Range** Routine Results;

N = Total number of PFUs counted on 4 the plates,

The maximum readable limit on PFUs count is 1000 on the four plate,

PFU count exceeding 1,000/4 plate is considered TNTC or >1,000,000 PFU/100gm

$$\text{Result} = \frac{(N \text{ PFUs}) * 100}{.1 \text{ ml}} = N * 1,000 \text{ PFU/100ml}$$

Example: High range version plate counts - 13, 23, 12, and 16 PFUs

$$\text{Result} = (64)*(1000) = 64,000 \text{ PFU/100ml}$$

Low Range Routine:

For each low range (effluent) sample, eight (8) Bottom Agar plates and eight (8) 2.5 ml DS Soft Agar tubes are prepared. Always begin the day's analyses with a negative control (blank) plate and finish analyses with a positive control plate and a negative control plate.

1. Allow prepared samples (50ml conical tubes, racked and labeled) to warm to room temperature immediately before analysis (20-30 minutes)
2. Remove tubes containing appropriate timed host cells from incubator immediately before analysis.
3. Vortex sample in 50ml conical tube for 10 seconds.
4. Moving quickly and smoothly, gently pipette 200µL of host cells into each of 8 soft agar tubes using a 1 ml serological pipet or 200µL micropipette with sterile tip.
5. Immediately thereafter, pipette 2500µL aliquot of sample into each of the 8 soft tubes using a 10 ml serological pipet and pipet aid or 2500µL micropipette with sterile tip.

6. Mix sample, host and soft agar by gently rolling test tube between hands.

Note: *Once E. coli F_{amp} is added to the mixture do not shake, only gently mix contents by rolling the tube between palms.*

7. Then overlay the contents of the soft agar tubes onto a Bottom Agar plates. Drag the mixture into a clear area and gently swirl the plates to spread sample and agar mixture.
8. Allow plates to set then inverted and incubated for 16 - 20 hours at 35- 37°C.
9. Circular zones of clearing (typically 1 to 10 mm in diameter) in lawn of host bacteria after 16 – 20 hours of incubation are counted as plaques. PFUs on each plate are counted and totaled using a digital counter pen and adequate light
10. Calculations of **Low Range** Routine Results;

N = Total number of Plaque forming units (PFUs) counted on 8 the plates,

The maximum readable limit on PFUs count is 2000 on the eight plate,

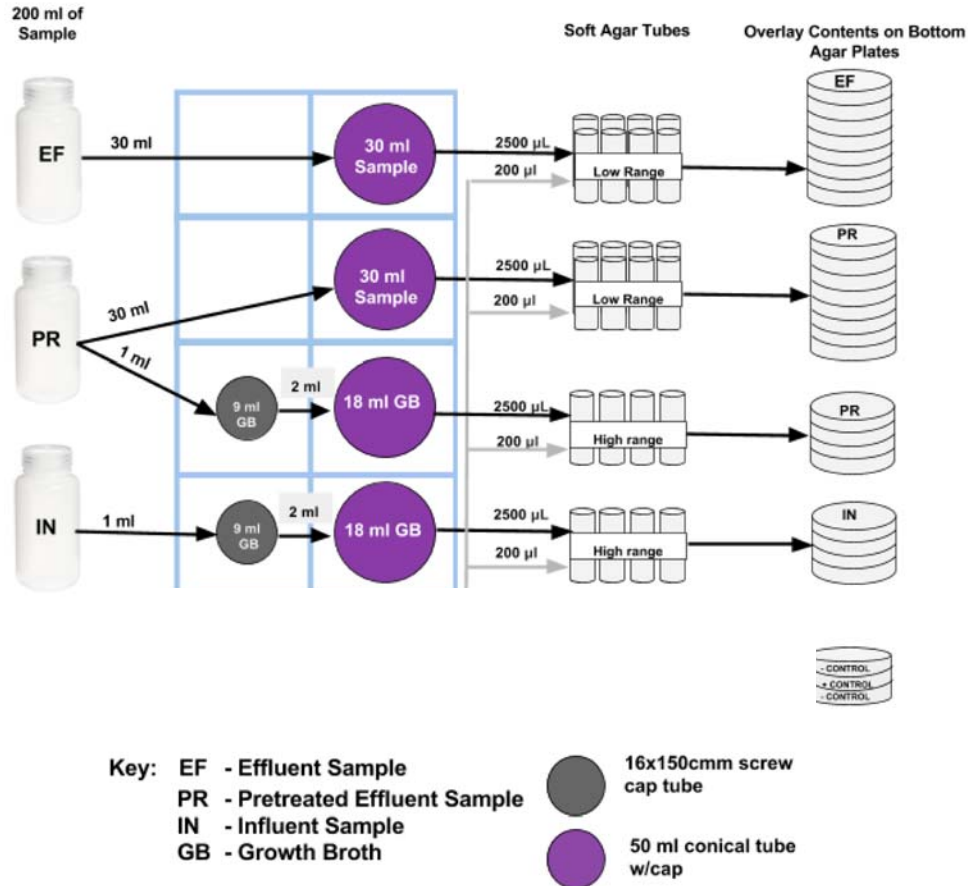
PFU count exceeding 2,000/8 plates is considered TNTC or >10,000 PFU/100gm

$$\text{Result} = \frac{(N \text{ PFUs}) * 100}{20 \text{ ml}} = N * 5 \text{ PFU}/100\text{ml}$$

Example: High range version plate counts - 21, 17, 20, 19, 13, 23, 12, and 16 PFUs

Result = (141)(5) = 702 PFU/100ml. Rounding off result to nearest 10s... Result = 700PFU/100gm*

MSC Method for Wastewater Schematic:



Samples Bench Sheet:

<p><i>Sample Bench Sheet</i> For use with the NSSP Method for Determination of MSC in Wastewater 2016 Edition</p>												
Initiation Analysts: <u>Jerry Jone</u>				Date: <u>10/10/16</u>				Time: <u>1:00PM</u>				
Completing Analysts: <u>Bill Bellicheck</u>				Date: <u>10/11/16</u>				Time: <u>8:00AM</u>				
Low Range Routine												
Sample#:1EF	WWTP: Hopetown, NE			Type: Effluent			Date/Time: 10/9/16					
PFU Counts	3	2	0	1	0	2	0	1	=	9	45 PFU/100ml	
Sample#:3Pre	WWTP: Hopetown, NE			Type: Pre-treatment			Date/Time: 10/9/16					
PFU Counts	199	215	203	170	233	210	206	188	=	1,624	8,120 PFU/100ml	
Sample#:	WWTP:			Type:			Date/Time:					
PFU Counts									=			
High Range Routine												
Sample#:2In	WWTP: Hopetown, NE			Type: Influent			Date/Time: 10/9/16					
PFU Counts	171	193	201	177	=	742					177,000 PFU/100ml	
Sample#:	WWTP: Hopetown, NE			Type: Pre-treatment			Date/Time: 10/9/16					
PFU Counts	3	1	2	0	=	6					6,000 PFU/100ml	
Sample#:	WWTP:			Type:			Date/Time:					
PFU Counts					=							
For unknown or mid-range sample, run both routines above on the sample												
Controls:	Start Neg Control:	count		0	End Pos Control:	count		43	End Neg Control:	count		0

F. Sample Collection and Storage.

1. Record all pertinent information on the collection form.
2. During transportation store samples in a cooler at 0 to 10°C
3. At laboratory, store samples in a refrigerator at 0 to 4 °C.
4. Maximum holding times for wastewater samples is up to 72 hours.

G. Quality Assurance.

1. Positive and negative control plates are run with MSC analyses each day.
2. Media sterility checks are made per batch and records are maintained.
3. Media log book is maintained (pH, volume, weights of each components, lot numbers, etc.).
4. An intra- and inter-laboratory performance program is developed.
5. Circular zones of clearing (typically 1 to 10 mm in diameter) in lawn of host bacteria after 16- 20 hours of incubation are counted as plaques. (Count the number of plaques on each plate.)
6. MSC determinations are reported as plaque forming unit (PFU) per 100 grams.
7. The desired range for counting is 0 to 300 PFU per plate. If the count exceeds the upper range or if the plaques are not discrete, results should be recorded as “too numerous to count” (TNTC) or >12,000 PFU/100ml for **Low Range** Routine and >1,200,000 PFU/100ml for **High Range** Routine.
8. Temperatures incubators are checked twice daily (at least 4 hours apart) to ensure operation within the stated limits of the method, and results are recorded in a logbook.
9. Check thermometers at least annually against a NIST-certified thermometer.
10. Calibrate the balance monthly using ASTM-certified Class 1 or 2 or NIST Class S reference weights.
11. Laboratory analysts adhere to all applicable quality control requirements set forth in the most recent version of FDA's *Shellfish Laboratory Evaluation Checklist*.

12. Calibration of micropipettes needs to be checked quarterly and records kept. Micropipettes used for handling MSC control and transferring host cells need to have a barrier tip or be dedicated to the specific use to prevent contamination

H. Safety.

Samples, reference materials, and equipment known or suspected to have sewage, coliphage and/or *E.coli* attached or contained must be sterilized prior to disposal.

I. Technical Terms.

- °C - degrees Celsius
- µL - microliter
- g - gram
- L - liter
- M - molar
- ml - milliliter
- Ave. - average
- MSC - Male-specific Coliphage, Male-specific Bacteriophage, F+ Bacteriophage
- NIST - National Institute of Standards and Technology
- PFU - plaque forming units
- RT - room temperature
- TNTC - too numerous to count

Host Strain: *E.coli* F_{amp} bacteria (*E.coli* HS(pFamp)RR)

Male-specific Coliphage: Viruses that infect coliform bacteria only via the F-pili.

Plaque : Clear circular zones (typically 1 to 10 mm in diameter) in lawn of host cells after incubation.

References:

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Pouillot, R., J.M. van Doren, , J. Woods, , D. Plante, M. Smith, G. Goblick, C. Roberts, A. Locas, W. Hajen, J. Stobo, J. White, J. Holtzman, E. Buenaventura, W. Burkhardt III, A. Catford, R. Edwards, A. DePaola, and K.R. Calci, 2015. Meta-analysis of the reduction of norovirus and male-specific coliphage concentrations in wastewater treatment plants. Appl. Environ. Microbiol. 81, 4669-4681.

U.S. Food and Drug Administration. 2004. Male-specific Coliphage (MSC) Workshop, conducted in Gloucester, Massachusetts on March 9-12, 2004.

Other Information:

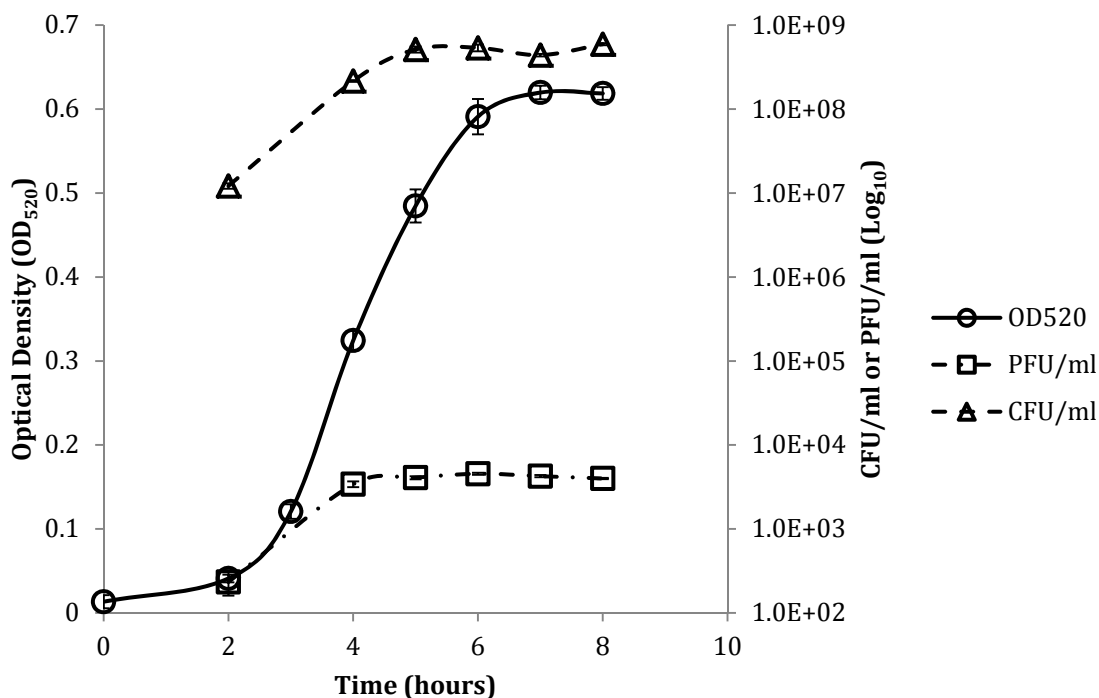
This method for the enumeration of male-specific coliphage in wastewater samples is inexpensive, easy to perform, and rapid, providing results within 24 hours. The cost of laboratory glassware, plastic-ware, agars, and reagents is approximately \$18 per series of samples (an influent, a pre-disinfection effluent, and a final effluent sample). In a well set-up laboratory, the method requires 6 hours of time from initiating host to pouring plates. Hands on technician time to perform this test is significantly less on the order of 1-4 hours per test depending upon how many tests are done per day. There are no special skill sets required beyond those required to operate a state-approved shellfish laboratory under the NSSP.

To standardize these assessments, an index of viral performance for use in the NSSP to determine classification options adjacent to WWTP outfall can be estimated. Subtracting the log value of final effluent score from the log value of the corresponding raw influent score will yield an index of viral performance ranging from 0 to 5. A viral index of 4 to 5 indicates 99.99% to 99.999% reduction of enteric viruses and would be considered high performance. A viral index of 1 to 2 indicates 90% to 95% reduction of enteric viruses through the treatment process and would be considered poor performance. A viral index <1 would be considered ineffective and should lead the SSCA to consider 10,000:1 rather than 1000:1 for the determination of the size of the prohibited zone adjacent to the outfall.

C. Validation Criteria

Determination of Optimal Optical Density of Famp Host (OD)

Procedures for enumeration of double-agar overlay method for male-specific coliphage provide different ranges for OD of host growth. Effort was taken to determine the working range of the host *E. coli* at 520nm, which is the current EPA measurement. Graph 1 shows that at approximately 4 to 6 hours growth the OD₅₂₀ of host is in the range of 0.35 to 0.6, during which time the MS2 plaquing efficiency of the host *E. coli* is optimal and consistent. Therefore, we conclude that a host OD₅₂₀ of 0.35 to 0.7, or approximately 4 to 6 hours of growth, is ideal for MSC enumeration.



Graph 1. Optical Density (OD₅₂₀) of *E. coli* HS(pFamp)RR in tryptone broth compared to plaque forming units (PFU) of MS2 coliphage. 10 ml of tryptone broth was inoculated with *E. coli* and incubated at 35°C. OD was measured every hour starting at t = 2 hours. At these intervals 100 µL of host was serially diluted and 100 µL of pre-determined dilutions were plated to determine CFU/mL. 200 µL of the same host sample was used to determine PFU/ml of stock MS2 controls.

The Determination of LOD, LOQ, and Linear Range using the NSSP SOP for the Single Laboratory Validation of Marine Biotxin and Non-MPN Based Microbiological Methods.

The SOP for the determination of LOQ, LOD, and the Linear Range is the most robust of the SOPs and yields a database from which subsets of data can be used to generate other validation criteria. For this database ten trials were run using clean effluent samples. Great effort was taken to find and verify clean effluent. The Dover, NH WWTP and the Hampton, NH WWTP were identified as high performing plants of different design capable of consistently producing clean effluent. Table 1 lists the metadata for the effluent samples collected for these trials

Table 1. Effluent Samples used for the Determination of LOD, LOQ, Linear Range

Trial #	Date Sampled	WWTP	Treatment Process
1	4/11/17	Dover	Tertiary, UV Dis-infection
2	4/11/17	Hampton	Secondary, Chlorine
3	4/18/17	Dover	Tertiary, UV Dis-infection
4	4/18/17	Hampton	Secondary, Chlorine
5	4/24/17	Dover	Tertiary, UV Dis-infection
6	4/24/17	Hampton	Secondary, Chlorine
7	5/1/17	Dover	Tertiary, UV Dis-infection
8	5/1/17	Hampton	Secondary, Chlorine
9	5/8/17	Dover	Tertiary, UV Dis-infection
10	5/8/17	Hampton	Secondary, Chlorine

For each of the 10 validation trials, 150 ml of clean effluent sample was aseptically transferred into 5-200ml sterile dilution bottles. A master spike solution was prepared in growth broth and was varied in concentration during the trials. The master spike solution was on the order of 10^3 MSC/ml. Four subsequent serial dilutions were made for each trial from the master spike at a 3:1 dilutions. This represented different spike concentrations over the working range of the method. The 5 dilution bottles were aseptically spiked with 5 ml of spike concentration 1 through 5, shaken vigorously and then 4 aliquots of 30 ml were transferred into 4-50 ml sterile conical tubes for each spike concentration (3 conical tubes for the replicates and a 4th tube for spike determination). In this way, 3 true replicates were generated at each of the 5 spike concentrations. This methodology was consistently applied throughout the ten trials. The 5 sets of 3 aliquots were processed and plated according to the method description above. Clean effluent was used instead of growth broth for the spike determination. Spike determinations using growth broth were underestimating the sample results. This problem was solved by using the same clean effluent similarly spiked. This strategy was employed as there is no standard method available for a truly independent spike determinations.

Table 2 below shows the Spiking Study Database for MSC Method in Wastewater SLV. Spike Concentrations and MSC replicate plate count results are in units of PFU of MSC/100ml.

Table 2. Spiking Study Database for MSC Method in Wastewater SLV

WW Spiking Database		8 plate				
Date	X Value	Replicate Plates	Log of plates	Y Value	Log RSD	X Value log of spike
	Measured Spike (PFU/100ml)			(PFU/100ml)		
	3400	4105	3.613	0.0025	-2.603	3.531
		4140	3.617			
		3980	3.600			
	1050	1025	3.011	0.0134	-1.874	3.021
		1225	3.088			
		1180	3.072			
Trial 1						
Dover	315	245	2.389	0.0352	-1.453	2.498
4/11/17		330	2.519			
		360	2.556			
	60	75	1.875	0.0629	-1.202	1.778
		60	1.778			
		45	1.653			
	10	30	1.477	0.2090	-0.680	1.000
		10	1.000			
		30	1.477			
	5950	5490	3.740	0.0046	-2.340	3.775
		5110	3.708			
		5155	3.712			
	1515	1355	3.132	0.0051	-2.289	3.180
		1365	3.135			
		1450	3.161			
Trial 2						
Hampton	410	225	2.352	0.0304	-1.517	2.613
4/11/17		225	2.352			
		170	2.230			
	70	65	1.813	0.0204	-1.690	1.845
		55	1.740			
		60	1.778			
	25	25	1.398	0.0321	-1.494	1.398
		30	1.477			
		25	1.398			

	5135	4315	3.635	0.0063	-2.199	3.711
		4800	3.681			
		4550	3.658			
	980	1085	3.035	0.0139	-1.856	2.991
		1005	3.002			
Trial 3		1220	3.086			
Dover						
4/18/17	405	385	2.585	0.0241	-1.618	2.607
		315	2.498			
		415	2.618			
	75	65	1.813	0.0379	-1.422	1.875
		90	1.954			
		80	1.903			
	25	20	1.301	0.1072	-0.970	1.398
		25	1.398			
		40	1.602			

	5175	4925	3.692	0.0065	-2.189	3.714
		5300	3.724			
		5490	3.740			
	1130	1280	3.107	0.0103	-1.986	3.053
		1160	3.064			
Trial 4		1340	3.127			
Hampton						
4/18/17	355	280	2.447	0.0317	-1.499	2.550
		335	2.525			
		405	2.607			
	40	60	1.778	0.0590	-1.229	1.602
		100	2.000			
		75	1.875			
	20	25	1.398	0.0634	-1.198	1.653
		30	1.477			
		20	1.301			

	11575	10655	4.028	0.0128	-1.891	4.064
		12800	4.107			
		10220	4.009			
	2080	2025	3.306	0.0212	-1.674	3.318
		2650	3.423			
		2735	3.437			
Trial 5						
Dover						
4/24/17	525	680	2.833	0.0360	-1.444	2.720
		705	2.848			
		465	2.667			
	190	205	2.312	0.0179	-1.746	2.279
		185	2.267			
		170	2.230			
	20	45	1.653	0.2069	-0.684	1.301
		60	1.778			
		15	1.176			

	12210	11140	4.047	0.0076	-2.121	4.087
		12165	4.085			
		10580	4.024			
	2555	2720	3.435	0.0058	-2.239	3.407
		2510	3.400			
		2520	3.401			
Trial 6						
Hampton	495	555	2.744	0.0395	-1.403	2.695
4/24/17		350	2.544			
		395	2.597			
	85	90	1.954	0.1082	-0.966	1.929
		110	2.041			
		45	1.653			
	20	20	1.301	0.1396	-0.855	1.301
		35	1.544			
		15	1.176			

	4430	3530	3.548	0.0131	-1.882	3.646
		4370	3.640			
		4075	3.610			
	1035	1100	3.041	0.0256	-1.592	3.015
		780	2.892			
		880	2.944			
Trial 7	240	275	2.439	0.0188	-1.727	2.380
Dover		230	2.362			
5/1/17		230	2.362			
	65	80	1.903	0.1126	-0.949	1.813
		75	1.875			
		35	1.544			
	10	20	1.301	0.1305	-0.884	1.000
		15	1.176			
		10	1.000			

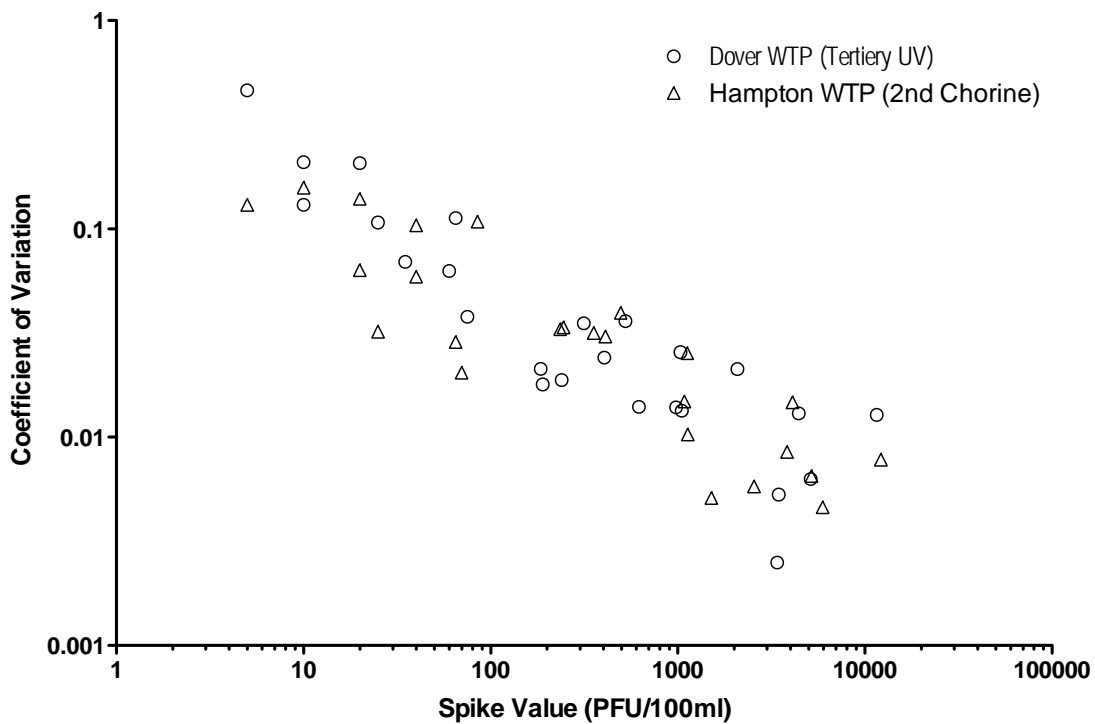
	4110	4415	3.645	0.0147	-1.833	3.614
		5630	3.751			
		5260	3.721			
	1125	955	2.980	0.0253	-1.596	3.051
		1060	3.025			
		1350	3.130			
Trial 8	245	315	2.498	0.0336	-1.474	2.389
Hampton		450	2.653			
5/1/17		325	2.512			
	40	35	1.544	0.1040	-0.983	1.602
		55	1.740			
		80	1.903			
	10	10	1.000	0.1580	-0.801	1.000
		10	1.000			
		20	1.301			

	3460	2765	3.442	0.0053	-2.273	3.539
		2940	3.468			
		3000	3.477			
	620	605	2.782	0.0140	-1.853	2.792
		725	2.860			
		650	2.813			
Trial 9						
Dover	185	210	2.322	0.0213	-1.672	2.267
5/8/17		175	2.243			
		215	2.332			
	35	35	1.544	0.0694	-1.158	1.544
		25	1.398			
		40	1.602			
	5	10	1.000	0.3618	-0.442	0.699
		5	0.699			
		3	0.477			

	3840	3490	3.543	0.0085	-2.073	3.584
		3675	3.565			
		4005	3.603			
	1085	825	2.916	0.0148	-1.831	3.035
		710	2.851			
Trial 10		855	2.932			
Hampton						
5/8/17	235	175	2.243	0.0330	-1.482	2.371
		135	2.130			
		185	2.267			
	65	60	1.778	0.0386	-1.413	1.813
		55	1.740			
		75	1.875			
	5	10	1.000	0.1305	-0.884	0.699
		15	1.176			
		20	1.301			

The replicate plate count results were log transformed and the relative standard deviation (RSD) were calculated. The RSD or coefficient of variation was plotted against the spike concentration and appears in Graph 2 below.

Graph 2. Coefficient of Variation verses Spike Concentration for Clean Effluent



To accurately determine the LOD and LOQ graphically, it was necessary to take the Coefficient of Variation and the Spike Determinations and to re-plot these as log values. Graph 3 below show the linear regression of the log transformed replicate and spiking data. Graphically, the LOQ/sensitivity of the method may be found at the point of intersection of the log spike concentration and the log coefficient of variation of -1.0 (or its antilog, 10%). The LOD may be found at the point of intersection of the log spike concentration and the log coefficient of variation of -0.477 (or its antilog of, 33%). Taking the antilog of the spike concentrations at these points of intersection gives the LOQ and LOD, respectively. Graph 3 indicates the LOQ and LOD for clean effluent to be 5.8 PFU/100gm and 0.9 PFU/100ml, respectively. The biostatistics program Prism 5.0 for Mac OS was used linear regression analysis and plots. The statistical summary of the linear regression from the log coefficient of variation verses log spike for the clean effluent data is presented in Table 3.

Graph 3. The LOD and LOQ/Sensitivity for Clean Effluent Samples

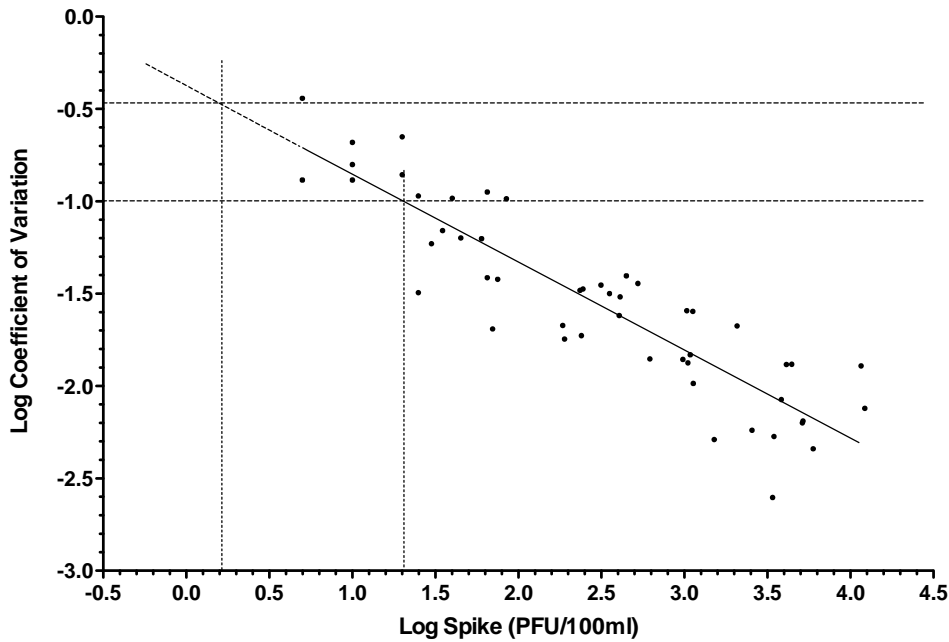


Table 3. Linear Regression Statistics for the Effluent Spiking Trials

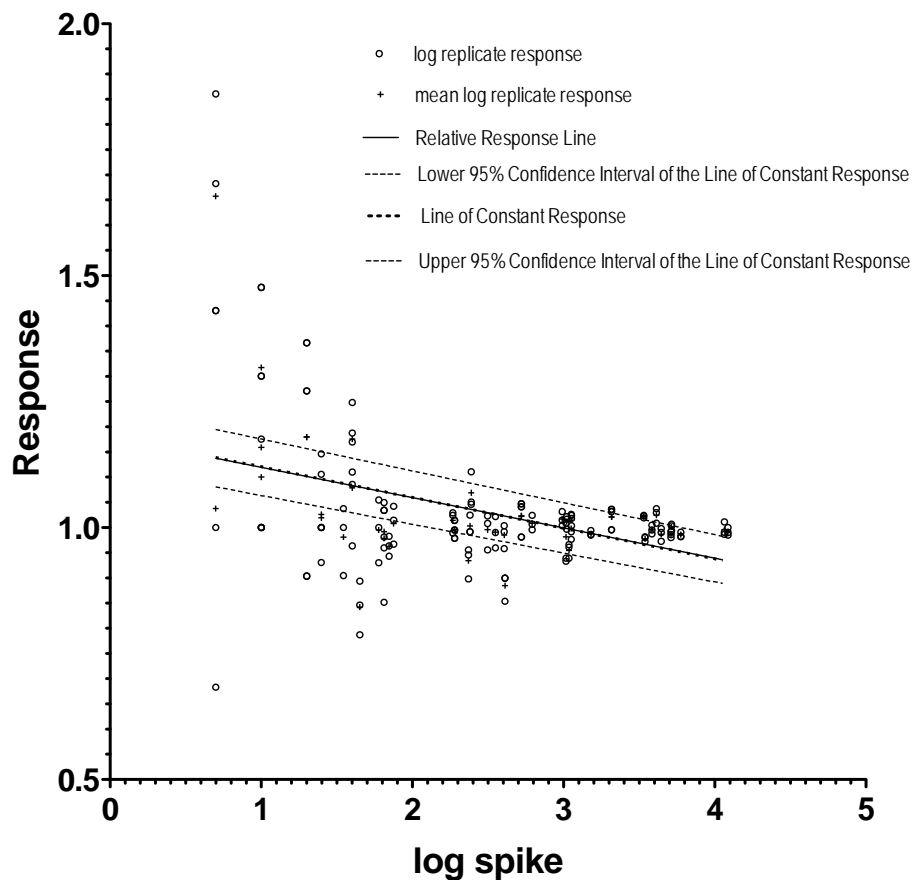
Best-fit values	
Slope	-0.4767 ± 0.03387
Y-intercept when X=0.0	-0.3746 ± 0.08890
X-intercept when Y=0.0	-0.7859
1/slope	-2.098
95% Confidence Intervals	
Slope	-0.5449 to -0.4085
Y-intercept when X=0.0	-0.5536 to -0.1957
X-intercept when Y=0.0	-1.345 to -0.3618
Goodness of Fit	
R square	0.8049
Sy.x	0.2244
Is slope significantly non-zero?	
F	198.1
DFn, DFd	1.000, 48.00
P value	< 0.0001
Deviation from zero?	Significant
Data	
Number of X values	50
Maximum number of Y replicates	1
Total number of values	50
Number of missing values	0

LOQ = Antilog [-2.098 (-1.0 + 0.375)] = 20.42 PFU/100ml
LOD = Antilog [-2.098 (-0.477 + 0.375)] = 1.63 PFU/100ml

The correlation coefficient (R square value) of this linear regression is 0.8049 which is above the threshold level of 0.64 and indicates a good fit. The LOD and LOQ as determined by the spiking trials shows LOD and LOQ of 1.63 PFU/100ml and 20.42 PFU/100ml, respectively. Rounding up, the LOD and LOQ are 2 PFU/100ml and 21 PFU/100ml, respectively.

To determine the **Linear Range**, data from Table 2 was manipulated to construct the relative response line, the line of constant response and the upper and lower 95% confidence interval bracketing the line of constant response as instructed in the SOP. Graph 4 below show that the upper (1.05) and the lower (.95) 95% confidence interval estimates are essentially parallel to the Relative Response line. This suggests that the method is linear through the working range of 5 PFU/100ml to 12,000 PFU/100ml.

Graph 4. Linear Range Determination



Data Summary:

Linear range of the method as implemented 5 to 12,000 PFU/100ml

The limit of detection of the method as implemented 2 PFU/100ml

The limit of quantitation/sensitivity of the method as implemented 21 PFU/100ml

Indeterminates <2 PFU/100ml to >12,000 PFU/100ml

The Determination of Accuracy/Trueness and Measurement Uncertainty is based upon the NSSP SOP for the Single Laboratory Validation of Marine Biotoxin and Non-MPN Based Microbiological Methods using the more robust databases acquired from the determination of the LOQ/LOD/Linear Range. The Accuracy/Trueness is calculated by dividing the log average of the plates by the log average of the spike concentrations, then multiplying the result by 100 to get a percent value. Table 4 shows the results for the Accuracy/Trueness of the method.

Table 4. Calculation of the Accuracy/Trueness of the MSC Method for Wastewater (Low Range Routine).

$$\begin{aligned} &\text{Average log of plates (2.473 PFU/100ml)/Average log of spike (2.455} \\ &\text{PFU/100ml)} \\ &= \text{Accuracy/Trueness of 100.7 \%} \end{aligned}$$

The Measurement Uncertainty is determined by subtracting the log mean replicate plate values from the reference or log spike values, then calculating the 95% confidence limits of the mean difference. Table 5 show the results of statistical analysis for Method Uncertainty.

Table 5 – Measurement Uncertainty in wastewater using low range routine.

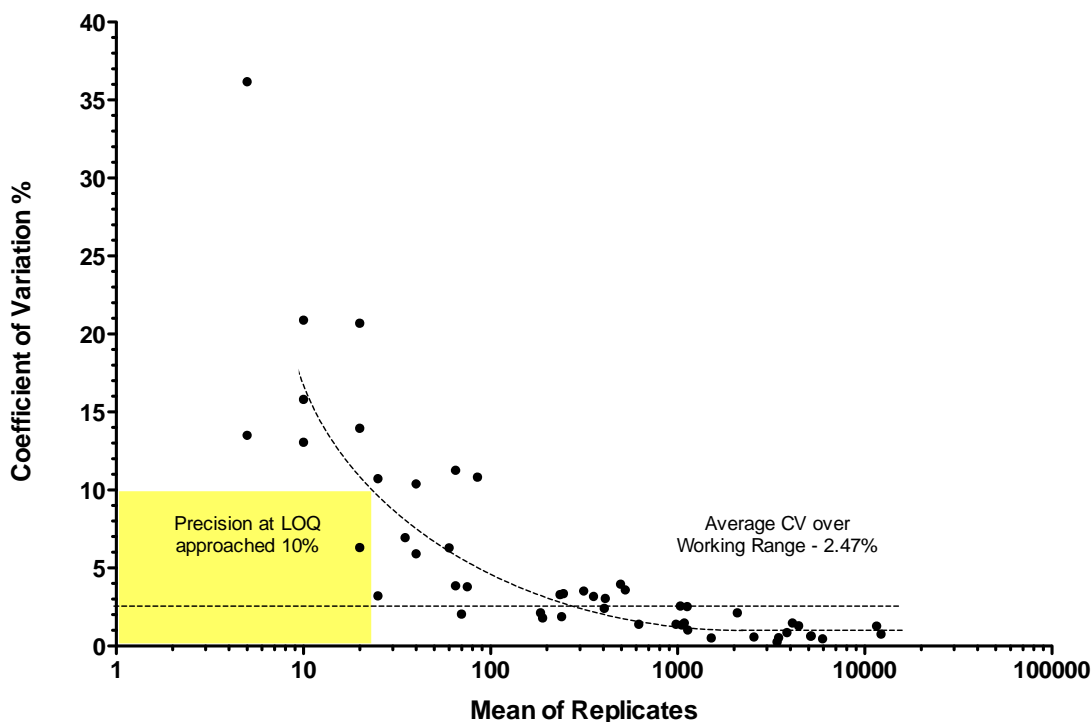
		Antilog
Number of values	150	
Mean	-0.01787	0.960
Std. Deviation	0.1566	
Std. Error	0.01278	
Lower 95% CI of mean	-0.04314	0.905
Upper 95% CI of mean	0.00739	1.017

Data Summary: Wastewater using Low Range Routine

Calculated % accuracy/trueness 100.7 %
 Calculated measurement uncertainty 0.905 to 1.017

The Determination of the Precision and Recovery is based upon the NSSP SOP for the Single Laboratory Validation of Marine Biotoxin and Non-MPN Based Microbiological Methods using the more robust data set acquired from the determination of the LOQ/LOD/Linear Range. To examine the precision over the working range of the method, a simple graphical approach was followed. The coefficients of variation were determined from the log transformed replicate data (50 sets of three true replicates) and were plotted verses the mean of the triplicate results (non log transformed data). The results are shown in Graph 5 for effluent using the low range routine.

Graph 5 - Coefficient of Variability (%) of Replicate verses Mean of Replicate for Effluent Samples using the Low Range Routine of the Method.



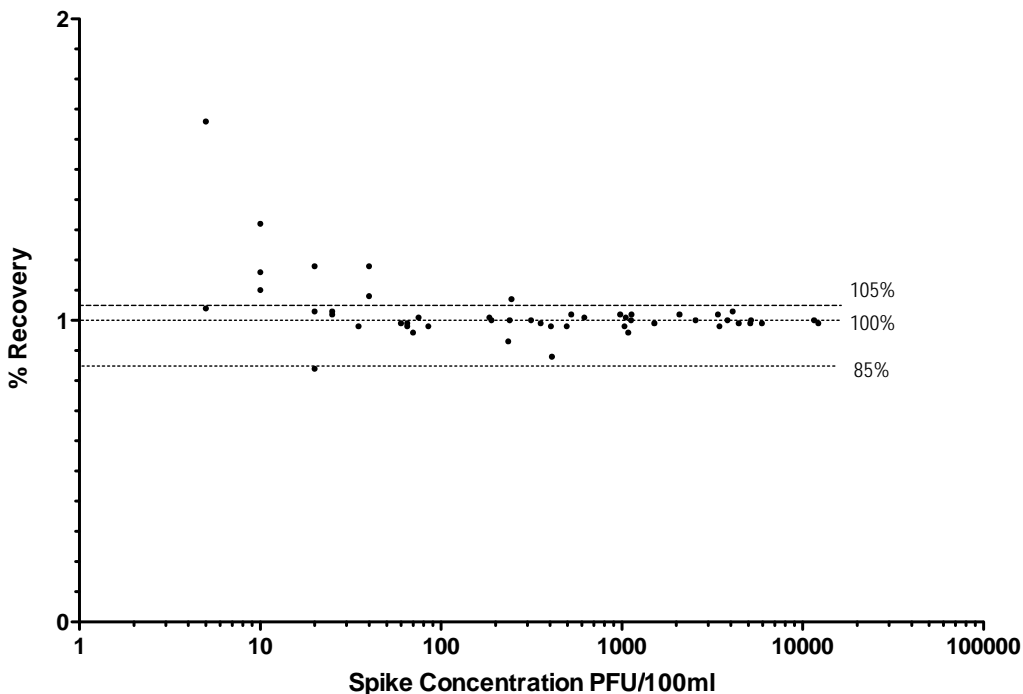
In Graph 4 above, the precision decreases as the LOQ and LOD are approached. The mean, minimum, and maximum coefficient of variations as determined over the working range for effluent samples appear in Table 6 below.

Table 6 – Mean, Minimum, and Maximum Coefficient of Variation over the Working Ranges.

Average Coefficient of Variation = 2.47%
 Minimum Coefficient of Variation = 0.25%
 Maximum Coefficient of Variation = 36.2%

To examine the **Recovery** over the working range of the method, a simple graphical approach was followed. The data from the LOD/LOQ/Linear Range was used for this determination. The mean of replicates was divided by the spike concentration. The percent recovery was then plotted against the spike concentrations. Graph 6 show these recovery plots with the recoveries bracketed at 85% and 105% for clean wastewater samples using the low range method routine. Recovery by the method is highly variable due to the problems associated with spike determinations. However, recovery for the method over all is high at 98.8%. (see Table 7)

Graph 6 - Percent Recovery verse the Spike Concentration for effluent using the low range method routine



As indicated above, the percent recovery of the method as implemented by this laboratory was calculated by dividing the log average of the replicates by the log spike concentration and multiplying by 100 to get a percent. Table 6 below shows this calculation from effluent trials for the low range method routine

Table 7 – Method Recovery from effluent trials.

Average	Average	%
Log of Spike	Log Replicates	Recovery
2.455	2.473	100.7%

Data Summary:

- Is the precision of the method under study consistent through the working range?
N, It varies as expected as the method approaches the LOD
- The coefficient of variation of the test method as implemented is **2.5%**
- Is the recovery of the method under study consistent through the working range?
N, It varies as expected as the method approaches the LOD
- What is the overall percent recovery of the method under study? **100.7%**

Ruggedness was determined using the NSSP SOP for the Single Laboratory Validation of Marine Biotxin and Non-MPN Based Microbiological Methods.

Two batches of bottom plates, soft agar tubes, and growth broths were prepared using two different lots of granulate agar (Media A and Media B), well in advance of the trials. Ten samples of clean effluent from the Dover and Hampton WTPs were similarly spiked and plated using media A and media B batches. The spike level was varied throughout the experiment. Table 8 shows the data, data analysis, and the results of the paired t-test for effluent

Table 8 - Determination of the Method Ruggedness for Effluent

<u>Media A</u> PFU/100gm	<u>Media B</u> PFU/100gm	Log Media A	Log Media B
3000	3575	3.48	3.55
5130	5055	3.71	3.70
5335	5465	3.73	3.74
14055	14980	4.15	4.18
13370	15955	4.13	4.20
12275	12200	4.09	4.09
5545	4845	3.74	3.69
5340	4495	3.73	3.65
210	235	2.32	2.37
190	175	2.28	2.24
	Skew	-1.35	-1.29
	Variance	0.47	0.48
	Ratio of Larger Var to Lower Var	0.01	

skew between -2 and 2 indicates symmetry

Ratio of Variances < 2 indicates homogeneity of variance

Paired t-test (Media A versus Media B)

P value	0.0.7648
P value summary	ns
Are means signif. different? (P < 0.05)	No
One- or two-tailed P value?	Two-tailed
t, df	t=0.3084 df=9
Number of pairs	10

Data Summary:

Value for the test of symmetry of the distribution of Media A data -1.35

Value for the test of symmetry of the distribution of Media B data -1.29

Variance of Media A data .047

Variance of Media B data .048

Ratio of the larger to the smaller of the variances of Media A and Media B 0.01

Is there a significant difference between Media A and Media N

Single Laboratory Validation (SLV) Protocol for Submission to the Interstate Shellfish Sanitation Conference (ISSC) For Method Approval

Name of the New Method:

Liquid Chromatography Tandem Mass Spectrometry (LC-MS/MS) Method for the Determination of Diarrhetic Shellfish Poisoning (DSP) Toxins in Shellfish.

Name of Method Developer:

Whitney Stutts, Ph.D. and Jonathan Deeds, Ph.D.

Developer Contact Information:

FDA Center for Food Safety and Applied Nutrition
5100 Paint Branch Parkway
College Park, MD 20740

Phone: 240-402-1474 (Deeds) or 240-402-1895 (Stutts)

whitney.stutts@fda.hhs.gov; jonathan.deeds@fda.hhs.gov

A. Need for the New Method

1. Need for Which the Method has been Developed:

Shellfish contaminated with natural toxins can cause consumer illnesses. The Food and Drug Administration has established guidance levels for five groups of natural toxins in shellfish responsible for the following illnesses: amnesic shellfish poisoning (ASP), azaspiracid shellfish poisoning (AZP), diarrhetic shellfish poisoning (DSP), neurotoxic shellfish poisoning (NSP), and paralytic shellfish poisoning (PSP). Shellfish hazards for domestic products are managed under the National Shellfish Sanitation Program (NSSP), but at present approved and approved limited use methods are only available for ASP, NSP, and PSP. Shellfish harvesting closures have been required due to DSP toxins in excess of the established regulatory guidance level of 16 µg OA eq./100 g shellfish on the Texas Gulf Coast since 2008, in the Puget Sound region since 2011, and in the New England region since 2015. This report describes the validation of an LC-MS/MS method for DSP toxins for use in the NSSP for the control of this hazard in clams. Once found to be fit for purpose for clams, the method will be fully validated for the additional matrices of mussel and oyster. Due to the immediate need for approved methods for this toxin group it was felt that submitting this proposal now, with the available full SLV data for clam, was important. Preliminary data is available for mussel and oyster such that the method can be used for these matrices in an approved limited use capacity.

2. Purpose and Intended Use of the Method: The intention is for this method to be an Approved Method for Biotoxin testing for DSP toxins under the NSSP (for clams) and that it should appear in Section IV. (Guidance Documents), Chapter II. (Growing Areas), Section .14 (Approved Laboratory Tests), Table 2 (Approved Methods for Marine Biotoxin Testing) under the new heading: Biotoxin Type: Diarrhetic Shellfish Poisoning (DSP), and the applications should be (1) Growing Area Survey and Classification and (2) Controlled Relaying with the sample type of Shellfish for both. Preliminary data is also provided for the additional matrices of mussel and oyster such that the method should be included in Table 4 (Approved Limited Use Methods for Biotoxin Testing) for these matrices while additional SLV data is generated.

3. Need for the New Method in the NSSP, Noting Any Relationships to Existing Methods:

The regulatory guidance level in the U.S. for diarrhetic shellfish poisoning toxins (DSTs) is 16 µg total (free plus esterified) toxins/100 g shellfish. The European Union (EU) recently adopted LC-MS/MS as the reference method for lipophilic shellfish toxins, which include DSTs.¹ However, because the EU harmonized protocol also measures a number of additional lipophilic toxins, including pectenotoxins, yessotoxins, and azapiracids, the protocol contains multiple method modifications and variations to account for this, depending on the needs of individual laboratories. This work optimized the EU lipophilic toxin reference method specifically for the quantitation of DSTs in clams for use in the NSSP. Some labs in the U.S. are already using best available science based on the EU LC-MS/MS reference method; thus, an NSSP-validated method for use in the U.S. is urgently needed. This LC-MS/MS method would be complimentary to other available testing methods such as the *in-vitro* protein phosphatase inhibition assay (PPIA), which has also been submitted for approval to the ISSC under a previous proposal, and comparative data is provided between these two methods in the comparability section.

4. Type of Method (Chemical, Molecular, or Culture):

Chemical Confirmatory Method: Liquid Chromatography Tandem Mass Spectrometry (LC-MS/MS) specifically measures okadaic acid, dinophysistoxin 1, and dinophysistoxin 2.

B. Method Documentation

Method Title:

Liquid Chromatography Tandem Mass Spectrometry (LC-MS/MS) Method for the Determination of Diarrhetic Shellfish Poisoning (DSP) Toxins in Shellfish.

Method Scope:

This method is fully validated for the determination of diarrhetic shellfish poisoning toxins in clams. Preliminary and comparative data only is provided for mussels and oysters.

References:

1. European Union Reference Laboratory for Marine Biotoxins. Harmonized Standard Operating Procedure for Determination of Lipophilic Marine Biotoxins in Molluscs by LC-MS/MS. Version 5, January 2015.
2. Gerssen, A.; McElhinney, M.A.; Mulder, P.P.J.; Bire, R.; de Boer, J. Liquid chromatograph-tandem mass spectrometry method for the detection of marine lyophilic toxins under alkaline conditions. *Journal of Chromatography A*, 1216 (2009) 1421-1430.
3. ICH Q2B, Validation of Analytical Procedures: Methodology, Fed. Regist. 1997, 62 (96), 27463-27467.
4. McNabb, P.; Selwood, A. I.; Holland, P. T. Multi-residue method for determination of algal toxins in shellfish: single-laboratory validation and inter-laboratory study. *Journal of AOAC International*, 88 (2005) 761.
5. AOAC International "Appendix K: Guidelines for Dietary Supplements and Botanicals, Part 1 AOAC Guidelines for Single-Laboratory Validation of Chemical Methods for Dietary Supplements and Botanicals", 2013.
6. Smienk et al. 2012 Single Laboratory Validation of A Ready-to-Use Phosphatase Inhibition Assay for Detection of Okadaic Acid Toxins. *Toxins*, 5, 339-352.
7. Smienk et al. 2013. Quantitative Determination of the Okadaic Acid Toxins Group by a Colorimetric Phosphatase Inhibition Assay: Interlaboratory Study. *Journal of AOAC International* Vol. 96, No. 1.

Principle:

Liquid chromatography tandem mass spectrometry is an analytical technique that provides both physical separation (liquid chromatography) and mass analysis (mass spectrometry) of sample components. In this work, a Waters ACQUITY® Ultra Performance Liquid Chromatography system was coupled with an AB Sciex 5500 QTrap® triple quadrupole mass spectrometer by electrospray ionization. Pre-validation studies were performed to assess the impacts of acidic versus basic chromatography and the use of neat versus matrix matched standard curves on overall method performance (data included). For targeted quantitation of DSTs, structurally informative transitions were specified for multiple reaction monitoring (MRM). In MRM mode, an ion of interest (precursor ion) is preselected in the first quadrupole and fragmented in the second quadrupole. Multiple product ions resulting from fragmentation of the precursor ion are then mass analyzed in the third quadrupole. Two analyte specific transitions, one for quantitation and one for confirmation, are specified for each analyte and monitored in both the calibration standard solutions and in the extracted shellfish matrices. A linear fit is applied to the peak area data for the quantitation ion collected for the calibration standards, and the equation for this line is utilized to calculate the concentration of each analyte in the spiked matrix samples. The enhanced resolving power afforded by LC and the selectivity gained by tandem mass spectrometry permitted the accurate detection and quantitation of DSTs in complex shellfish matrices. Individual toxin values are converted to a single integrated okadaic acid equivalents value through the use of established

toxicity equivalency factors (OA: 1, DTX1: 1, DTX2: 0.6). Only OA and DTX1 have been found to date in the U.S.

Analytes of Interest:

Diarrhetic shellfish poisoning toxins: Okadaic Acid (OA), Dinophysistoxin-1 (DTX1), and Dinophysistoxin-2 (DTX2). Shellfish metabolites (fatty acid acyl-esters for all 3 toxins, collectively referred to as DTX3) are included through the use of a required alkaline hydrolysis step.

Proprietary Aspects: None

Cost of the Method:

Capital equipment purchases:

1. Ultra Performance Liquid chromatograph (UPLC) or High Performance Liquid Chromatograph (HPLC), capable of running in gradient mode [example: Waters Corporation Acquity UPLC system (Manchester, UK) (approx. \$60,000, new with government (GSA) discount)]
2. Mass Spectrometer, equipped with an electrospray ionization source and multiple reaction monitoring scan mode capabilities [example: AB Sciex QTrap 5500 equipped with a Turbo V ionization source (Framingham, MA) (approx. \$320,000, new with government (GSA) discount)]

Cost per sample (approx. for consumables): \$10/sample

Sample Turn Around Time: Analysis of 10 samples, including extraction, hydrolysis, sample analysis, and quantitation can be accomplished in approx. 6 -7 hours. Analysis time increases by 2.5 hours for every 10 additional samples.

Equipment Required:

1. Instrumentation for sample preparation: knives for shucking, stainless steel laboratory spatulas, sieve for draining
2. 4 oz. plastic screw top specimen cups (if a sample of unextracted homogenate is to be saved) (e.g., Covidien #17099)
3. Balance, accuracy to the nearest 0.01 g
4. Blender or food processor
5. Laboratory homogenizer (e.g., Polytron, Ultraturax, etc.) equipped with a generator appropriate for shellfish (e.g. 20 mm saw toothed)
6. Vortex mixer (either single or multi-tube)
7. Centrifuge, capable of 2000 x *g*, with adaptors for 50 mL and 15 mL centrifuge tubes
8. Heat block or water bath capable of maintaining 76 ± 2 °C
9. Volumetric flasks or graduated cylinders capable of accurately measuring 20, 100, 500, and 1000 mL

10. Disposable 50 mL centrifuge tubes
11. Disposable glass Pasteur pipettes with bulbs
12. 20 mL glass scintillation vials with coned polyethylene lined caps (if a sample of non-hydrolyzed methanolic extract is to be saved) (e.g., Wheaton #986560)
13. Disposable 16 × 100 mm glass screw cap tubes with phenolic PTFE lined caps (e.g., Fisher #14-959 35AA (tubes), Corning #9998-15 (caps))
14. 0.01, 0.05, 0.1, 0.5, and 1 mL positive displacement microdispensers or syringes (e.g., Drummond #3-000-510, 3-000-575, 3-000-590 or Hamilton #80530, 80630, 81330, 81365)
15. 1 mL disposable syringes (e.g., Becton Dickinson #309602)
16. PTFE syringe tip membrane filters, 13 mm, pore size 0.2 µm (e.g., Pall #PN4542)
17. LC-MS autosampler vials with pre-slit caps (e.g., Agilent #5182-0715)
18. Reversed Phase HPLC Column (e.g., Acquity UPLC BEH C18 1.7 µm particle size, 1.0 × 150 mm)
19. Ultra Performance Liquid chromatograph (UPLC) or High Performance Liquid Chromatograph (HPLC), capable of running in gradient mode
20. Mass Spectrometer, equipped with an electrospray ionization source and multiple reaction monitoring scan mode capabilities

Reagents Required:

1. Certified Reference Standards

Okadaic acid: e.g. (NRC-CRM-OAc) NRC-CNRC Institute for Marine Biosciences, Canada

Dinophysistoxin-1: e.g. (NRC-CRM-DTX1) NRC-CNRC Institute for Marine Biosciences, Canada

Dinophysistoxin-2: e.g. (NRC-CRM-DTX2) NRC-CNRC Institute for Marine Biosciences, Canada

2. Chemicals

Acetonitrile, HPLC Optima Grade

Methanol, HPLC Optima Grade

Water, HPLC Optima Grade

Ammonium formate (≥98% purity)

Formic Acid (≥98% purity)

Hydrochloric acid (37%)

Sodium hydroxide (≥98% purity)

Hexanes (Certified ACS, ≥98.5%)

Solution Preparation

1. Extraction solvent: 100% methanol

2. Hydrochloric Acid 2.5 M: Add 60 mL of water to a 100 mL volumetric flask or graduated cylinder. To the water, add 20 mL of hydrochloric acid and then dilute with water to 100 mL. Place in an appropriate glass jar for storage marked with the date of creation. This solution may be stored at room temperature and can be used for up to three months.

3. Sodium hydroxide 2.5 M: Dissolve 10 grams of sodium hydroxide in 75 mL of water in a glass beaker and add to a 100 mL volumetric flask or graduated cylinder. Make up to 100 mL with water and transfer to an appropriate storage vessel marked with the date of creation. This solution may be stored at room temperature and used for up to 3 months.

Matrix or Matrices of Interest:

Clam, mussel, and oyster. Full SLV validation data is provided here for clam but pre-validation and method comparability studies (data provided) indicate that the method will also be applicable to additional shellfish matrices such as mussel and oyster. Data for additional matrices will be provided upon approval of the method for clam.

Sample Collection, Preservation, and Storage requirements:

CRM-DTX1, CRM-DTX2, CRM-OA stock solutions (in MeOH in sealed ampules) were purchased from the National Research Council Canada and stored at -20 °C according to manufacturer’s recommendations.

For each bivalve type, animals were collected from four different growing areas (Table 1). Shellfish were shucked, rinsed and drained to remove salt water. Ten to twelve animals were combined and homogenized using a blender/food processor and/or a Polytron homogenizer. Homogenized tissues were stored in plastic screw top specimen containers at -20 °C until used.

Table 1. Sources of blank shellfish matrices for spiking experiments

Source	Clam (<i>Mercenaria mercenaria</i>)	Mussel (<i>Mytilus edulis</i>)	Oyster (<i>Crassostrea virginica</i>)
A	Provided by Maryland Department of Natural Resources	Provided by MD Department of Natural Resources (used for pre-validation studies)	Provided by Maryland Department of Natural Resources (used for pre-validation studies)
B	Purchased live directly from harvester in Virginia	Washington Department of Health (used for method comparability data)	Provided by Texas Department of State Health Services (used for method comparability data)
C	Purchased live from retail. Harvest location Northport, Maine	Not performed yet	Not performed yet
D	Provided by Florida Wildlife Research Institute. Collection location Cedar Key, Florida.	Not performed yet	Not performed yet

Safety Requirements:

Proper precautions should be taken to avoid inhalation of harmful reagents or contact with skin or eyes. Analyst should wear a lab coat, gloves and safety glasses when working with chemicals. Chemical reagents that are flammable and/or toxic should be used within a chemical fume hood to protect laboratory workers.

Other Information:

Technical Skills Required: General laboratory skills (i.e., ability to accurately pipette small volumes, etc.). Experience with operation and general maintenance of liquid chromatography mass spectrometry equipment is required.

Abbreviation and acronyms:

DTX1: Dinophysistoxin-1

DTX2: Dinophysistoxin-2

OA: Okadaic acid

LC-MS/MS: Liquid Chromatography Tandem Mass Spectrometry

Step by Step Test Procedure:

Shellfish Preparation:

1. Clean outside of the shellfish with fresh water.
2. Cut the adductor muscles to open and rinse the inside with fresh water to remove any debris.
3. Remove the meat from the shell and drain tissue using a sieve to remove salt water.
4. Combine 10–12 animals and homogenize using a blender/food processor or a laboratory homogenizer.

Extraction Procedure:

1. Accurately weigh $2.00 \text{ g} \pm 0.05 \text{ g}$ of tissue homogenate into a 50 mL disposable centrifuge tube.
2. Add 9.0 mL of methanol and mix thoroughly using a vortex mixer for 3 min.
3. Centrifuge at 2000 *g* for 10 min at approximately 20 °C.
4. Transfer the supernatant to a clean 20 mL graduated cylinder, volumetric flask, or glass scintillation vial.
5. Re-extract the residual tissue pellet with an additional 9.0 mL of methanol and homogenize using a laboratory stick homogenizer (e.g. Polytron, Ultraturax).
6. Centrifuge at 2000 *g* for 10 min at approximately 20 °C.
7. Transfer the supernatant to a 20 mL volumetric flask or graduated cylinder and combine with the first extract. Adjust total extract volume to 20 mL with 100% methanol.
8. Transfer the 20 mL of total extract back into the 20 mL scintillation vial for storage (if desired).

Hydrolysis:

Note: For this procedure, total DSP toxins (i.e. free toxin plus toxin fatty acid esters) must be quantified. To do this, all samples must go through an alkaline hydrolysis step prior to analysis.

Accurately transfer a 2 mL aliquot of the 20 mL methanolic extract to a 16 × 100 mm glass tube with a phenolic PTFE lined screw cap using a positive displacement microdispenser or syringe. Add 250 µL of 2.5 M NaOH to the 2 mL extract aliquot. Homogenize with a vortex mixer for 30 seconds and record the total weight of the tube. Make sure caps are securely fastened and heat the mixture at 76 °C for 40 minutes. [Note: 76 °C is above the boiling point for methanol, therefore sample loss will occur if tube caps do not fit well and are not securely fastened.] Dry the water from the outside of the tubes, allow the tubes to cool to room temperature (approx. 5-10 minutes), and then re-weigh each tube. If the weight has dropped by more than 0.1 g, replace lost weight using 100% MeOH. Finally, neutralize samples with 250 µL of 2.5 M HCL, mix by vortexing for 30 seconds.

Sample Clean-up:

Partitioning of the methanolic shellfish extracts with hexane is performed to remove nonpolar lipids such as triglycerides, which could contaminate the source region of the instrument and/or suppress the responses of the DSTs.⁴ Add 5 mL of hexanes directly to the 2.5 mL hydrolyzed methanolic extract. Mix by vortexing for 1 minute. Partition by centrifuging at 2,000 g for 10 min. Using a disposable glass pasture pipette, collect the hexanes (top layer) and discard to an appropriate waste container. Transfer approximately 1 mL of the methanolic extract (bottom layer) into a 1 mL disposable syringe equipped with a 0.2 µm PTFE syringe tip filter. Filter directly into an LC-MS certified glass vial and cap.

Quality Control Steps:

1. If available, a divert valve should be used to divert LC flow at the beginning and end of each chromatographic run. If a divert valve is not used, the ion source region and curtain plate will need to be cleaned between each batch (approximately every 24 hours) to maintain adequate sensitivity.
2. Use a new calibration curve each day of analyses. The calibration curve should be analyzed before and after each set of samples, and the data from both curves should be fitted with a line using least squares linear regression. Each calibration curve should be derived from at least six calibration points and the linear regression should yield a correlation coefficient (R^2) greater than or equal to 0.98. Analysts should also visually inspect the plot of the calibration to confirm linearity. If a calibration curve yields a correlation coefficient less than 0.98 or if non-linearity is visually observed, a new calibration curve should be prepared and samples should be reanalyzed.
3. The variation in the calibration curve slopes between the first and second set of calibration standards should not exceed 25%.
4. Reagent Blanks (methanol solvent) should be analyzed after the high calibration standard and periodically after fortified samples to insure that analyte carryover is not occurring; toxins

should not be detected above 10% of the lowest calibration point or should be below LOD. If carryover is observed, the LC gradient should be extended to allow for a longer wash at high organic (99% B).

5. Procedural Blanks (methanol carried through sample preparation process at the same time as the samples) should be analyzed before and after extracted samples.
6. One mid-scale calibration standard (e.g., 10 ng/mL) should be analyzed bracketing at least every 10 samples to assure that no retention time shifts (possibly due to column failure) or loss in signal intensity (due to fouling of the column or mass spectrometer) has occurred.
7. The retention time of analytes in all matrix solution should be within 3% that of the neat toxin standards.
8. Each chromatographic peak must be defined by at least 10 data points.
9. To confirm the presence of each DST, two mass transitions must be observed above the LOD. The transition that yields the highest signal-to-noise ratio(S/N) is used for quantitation, and the transition yielding the second highest S/N is used for confirmation. The S/N of the peak used for confirmation must be ≥ 3 .
10. Ion ratios between the quantitative and qualitative ion transitions should be within $\pm 20\%$ that of the relative ion abundance of the neat toxin standards.
11. Chromatographic separation must be sufficient for resolving OA and DTX2. Peak resolution (R_s) of OA/DTX2 should be calculated using the equation below and must be ≥ 1 for correct identification).

$$R_s = \frac{2x(RT_2 - RT_1)}{W_1 - W_2}$$

LC-MS/MS Method

Instrumentation Used for Validation: AB Sciex QTrap 5500 equipped with a Turbo V ionization source (Framingham, MA) and a Waters Corporation Acquity UPLC system (Manchester, UK).

LC Parameters:

UPLC column used for validation: Waters BEH C18 (1.7 μm , 1.0 mm \times 150 mm)

Column Temperature: 40 $^{\circ}\text{C}$

Autosampler Temperature: 10 $^{\circ}\text{C}$

Injection Volume: 5 μL

LC Gradient:

Mobile phase A: 2mM ammonium formate and 50 mM formic acid in 100% water.

Preparation of 1000 mL: dissolve 128 ± 10 mg ammonium formate in water and transfer into a 1000 mL volumetric flask; fill approximately half-way to the mark with water and add 1.9 mL of formic acid. Fill to mark with water.

Mobile phase B: 2mM ammonium formate and 50 mM formic acid in 95% acetonitrile/5% water.

Preparation of 1000 mL: dissolve 128 ± 10 mg ammonium formate in 48.1 mL water and transfer into a 1000 mL volumetric flask; fill approximately half-way to the mark with acetonitrile and add 1.9 mL of formic acid. Fill to mark with acetonitrile.

Weak needle* and strong needle* wash solvent composition matched that of mobile phase A and B, respectively (*specific to Waters Acquity UPLC).

For the gradient in Table 2, LC flow should be diverted to waste from time 0.0–3.5 min and from 9.0–15.0 min.

Table 2. LC Gradient

Time (min)	Flow Rate (mL/min)	%A	%B
0.0	0.120	50	50
2.0	0.120	50	50
6.0	0.120	30	70
8.0	0.120	1.0	99
10.0	0.120	1.0	99
10.5	0.120	50	50
15.0	0.120	50	50

Analyte retention times for this gradient and column can be seen in Figure 1.

MS Ion Source Parameters:

Turbo V ion source parameters were optimized in negative ionization mode for all analytes under the acidic chromatographic conditions listed above. These parameters will vary between different instrument platforms or ionization sources.

Table 3. Turbo V ion source parameters

Source Temperature	550 °C
Ion Spray Voltage	-4500 V
Curtain Gas	25 au
Gas 1	40 au
Gas 2	40 au

MRM Parameters:

Manual or automatic compound optimization must be performed by every laboratory to determine the optimal settings for the MRM parameters as these may vary between different instrument platforms. Ruggedness studies (detailed later in the document) found that compound re-optimization was required even for detector replacement with the same make and model. Instrument must be correctly calibrated for negative ionization mode. OA, DTX1, DTX2 should be monitored in negative ionization mode. The precursor and product ion mass-to-charge (m/z) should be confirmed by acquiring full scan MS1 and MS/MS spectra for each toxin. At least two product ions must be monitored for each toxin, one for quantitation and one for confirmation. Dwell times for transitions should be set such that at least 10 data points are acquired across each peak. Declustering Potential (DP), Entrance Potential (EP), Collision gas (CAD), Collision Energy (CE), and Collision Cell Exit Potential (CXP) should be optimized for each MRM transition monitored. Table 4 lists the compound-dependent parameters optimized for two different AB Sciex QTRAP 5500 detectors. For method validation, two confirmatory ions were monitored; however, for routine analysis monitoring the product ion at m/z 151 is not required.

Table 4. MRM Parameters used for Method Validation on an AB Sciex QTrap 5500 Mass Spectrometer

Compound	Polarity	Q1 (m/z)	Q3 (m/z)	Dwell Time (ms)	DP [†] (V)	EP (V)	CAD Gas	CE (V)	CXP (V)
OA	Negative	803.5	255.2	100	-110	-10	High	-70	-15
OA	Negative	803.5	113.1	100	-110	-10	High	-100	-19
OA*	Negative	803.5	151.1	100	-110	-10	High	-70	-15
DTX2	Negative	803.5	255.2	100	-110	-10	High	-70	-15
DTX2	Negative	803.5	113.1	100	-110	-10	High	-100	-19
DTX2*	Negative	803.5	151.1	100	-110	-10	High	-70	-15
DTX1	Negative	817.5	255.2	100	-110	-10	High	-70	-15
DTX1	Negative	817.5	113.1	100	-110	-10	High	-100	-19
DTX1*	Negative	817.5	151.1	100	-110	-10	High	-70	-15

*One additional confirmatory ion transition was monitored for method validation purposes.

† Compound re-optimization for ruggedness testing using a second AB Sciex QTRAP 5500 detector found that the declustering potential had to be changed to -5 V for optimum method performance.

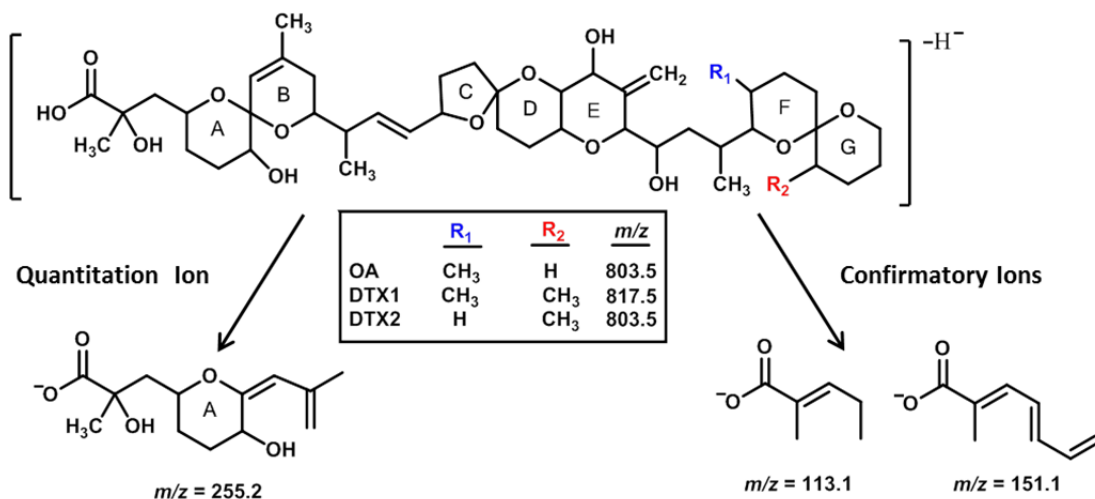


Figure 1. Structure of precursor ions and proposed product ion structures for DSTs.

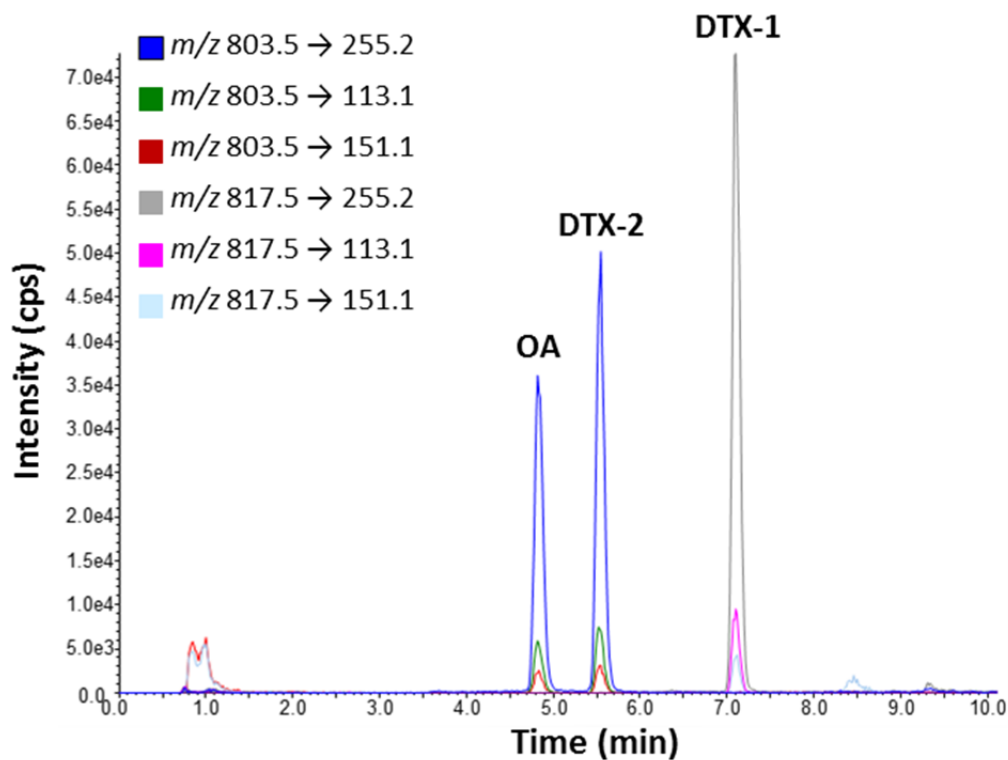


Figure 2. MRM chromatogram for the 12.5 ng/mL (regulatory level = 12.8 ng/mL) spike of OA, DTX1, and DTX2 into blank clam matrix.

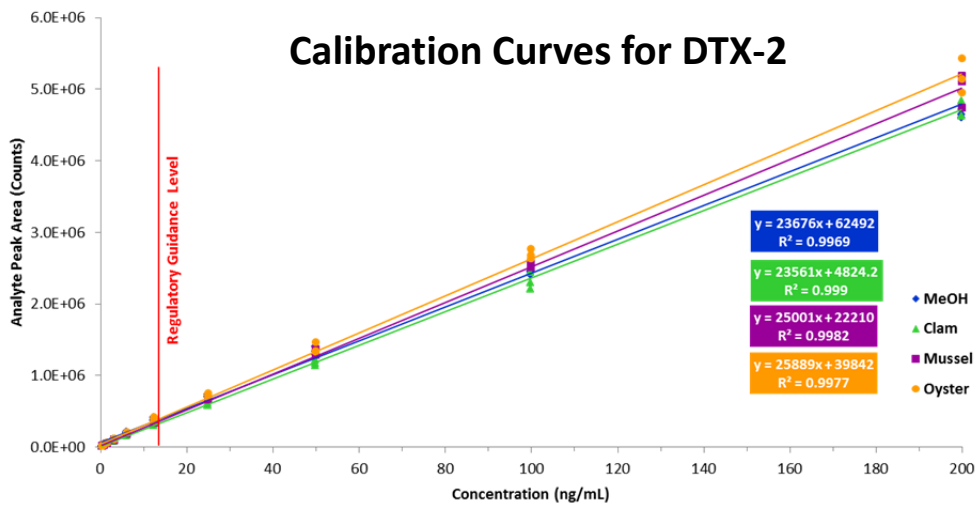
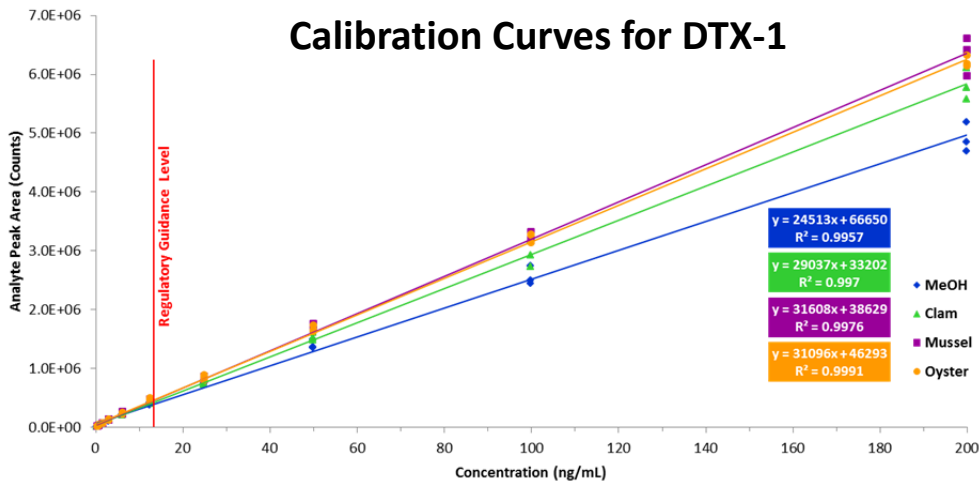
Pre-Validation Study Results (Testing for potential shellfish matrix effects for clam, mussel, and oyster, and ruggedness testing for acidic versus basic chromatography)

Initial pre-validation testing investigated the impact of mobile phase pH on method performance, and potential matrix effects (i.e. MS signal suppression or enhancement) for clam (*Mercinaria mercinaria*), mussel (*Mytilus edulus*), and oyster (*Crassostrea virginica*) in neat versus matrix matched spiked standard curves. Various laboratories in the United States and abroad are using different chromatographic mobile phase conditions, based on individual lab preference and need. Specifically, some labs are utilizing acidic chromatography (pH 2.4) while others are using basic chromatography (pH 11). Acidic chromatography is routinely used for quantifying total DSP toxin following hydrolysis. However, for methods that require polarity switching to investigate many classes of lipophilic toxins, basic chromatography is advantageous in that analytes that are preferentially ionized in positive mode are sufficiently separated from those that are preferentially ionized in negative mode.² Basic chromatography has also been reported to enhance sensitivity for certain lipophilic toxins, but due to matrix effects often requires the use of matrix matched standard curves, which adds additional cost and time to the method.

Figures 3 and 4 illustrate calibration curves from neat standard solutions in methanol compared to spiked methanolic extracts of clam, mussel, and oyster analyzed under acidic and basic chromatographic conditions, respectively. For the preparation of blank shellfish extracts, homogenates from 5 previously tested composite shellfish samples (10-12 animals each) for each species, found to have no detectable DSTs, were re-extracted, hydrolyzed, and hexane washed following the procedure described above. These 5 extracts were then pooled per species and used for the preparation of matrix-matched standard curves. Ten fortification concentrations: 0.391, 0.781, 1.56, 3.13, 6.25, 12.5, 25, 50, 100, and 200 ng/mL (equivalent to 0.49-250 µg/100g) of each toxin (from 2 µg/mL stocks) were used to construct each calibration curve. Fortification with DSTs was done post extraction to evaluate potential matrix interferences without having to account for analyte recovery. Each curve was analyzed in triplicate. All calibration curves were shown to be linear ($R^2 \geq 0.99$) within the range of 0.391–200 ng/mL using both acidic and basic chromatography.

Our results indicate that under acidic chromatographic conditions over an extended working range, the slope of the calibration curves for OA and DTX2 in the presence of matrix is within 10% that of the neat toxin standards in methanol for all three matrices. DTX1 response, on the other hand, showed signal enhancement in matrix compared to neat standards, especially at high toxin concentrations (>50 ng/mL in the hydrolyzed shellfish extract). The percent difference in the slope of the solvent-only calibration curve and the matrix-matched calibration curve for DTX1 was 19% in clam, 29% in mussel, and 27% in oyster. However, at the regulatory guidance level (16 µg/100g shellfish, 12.8 ng/mL on column, in the hydrolyzed shellfish extract), the difference in peak areas for DTX1 in matrix versus solvent is <20% for all three matrices. In comparison, under basic chromatographic conditions OA and DTX2 signals were suppressed in the presence of matrix, resulting in significantly lower calibration curve slopes (up to 19% lower) for matrix-matched calibration curves. At the regulatory guidance level, matrix suppression resulted in response differences as high as 40% for OA and 37% for DTX2. LODs and LOQs were comparable for both acidic and basic chromatography, and both were well below the level of concern for these toxins (additional information below). Based on this information, acidic chromatographic conditions were chosen for the SLV study because they would allow the use of neat standard curves in methanol, saving both time and expense. In addition, the greatest risk from the use of acidic chromatography with non-matrix-matched standard curves is potentially overestimating DTX1 concentrations (increased chance of false positive result of exceeding guidance level), while the greatest

risk from basic chromatography with non-matrix matched standard curves is underestimating OA or DTX2 concentrations (increased chance of false negative result of exceeding guidance level). From a public health perspective, overestimation is more protective than underestimation and it was felt that this slight risk was acceptable when weighted against the additional burden in both time and expense in requiring the use of matrix-matched standard curves. Furthermore, additional validation including the use of matrix matched standard curves from multiple matrix sources would likely be required for accurate quantitation of DSTs when using mobile phases at high (basic) pH.



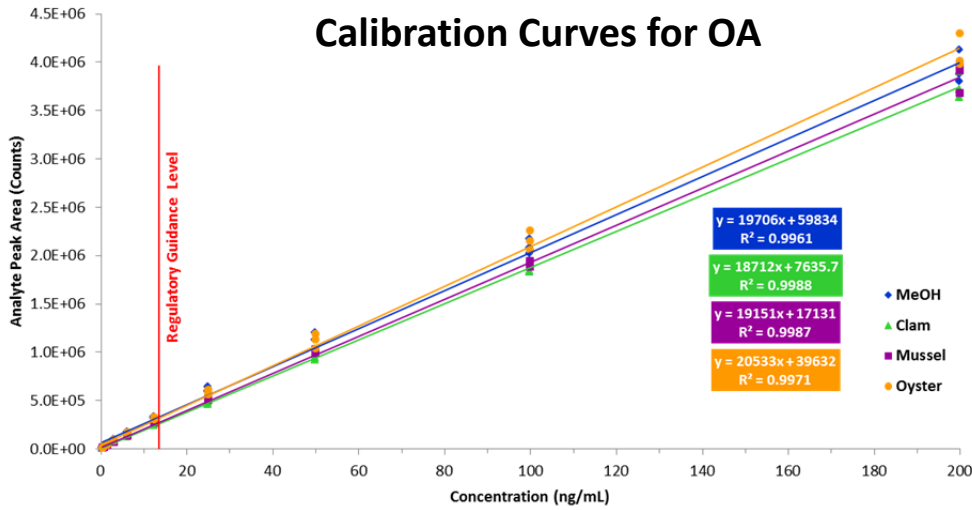
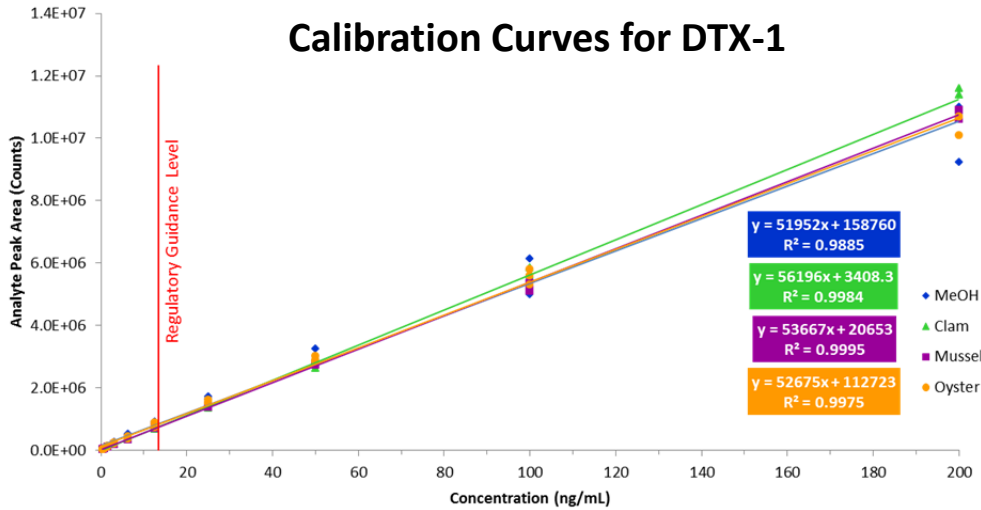


Figure 3. Calibration curves from neat standard solutions and spiked methanolic extracts of clam, mussel, and oyster analyzed under acidic chromatographic conditions. A working range of 0.39–200 ng/mL (n=10) was used.



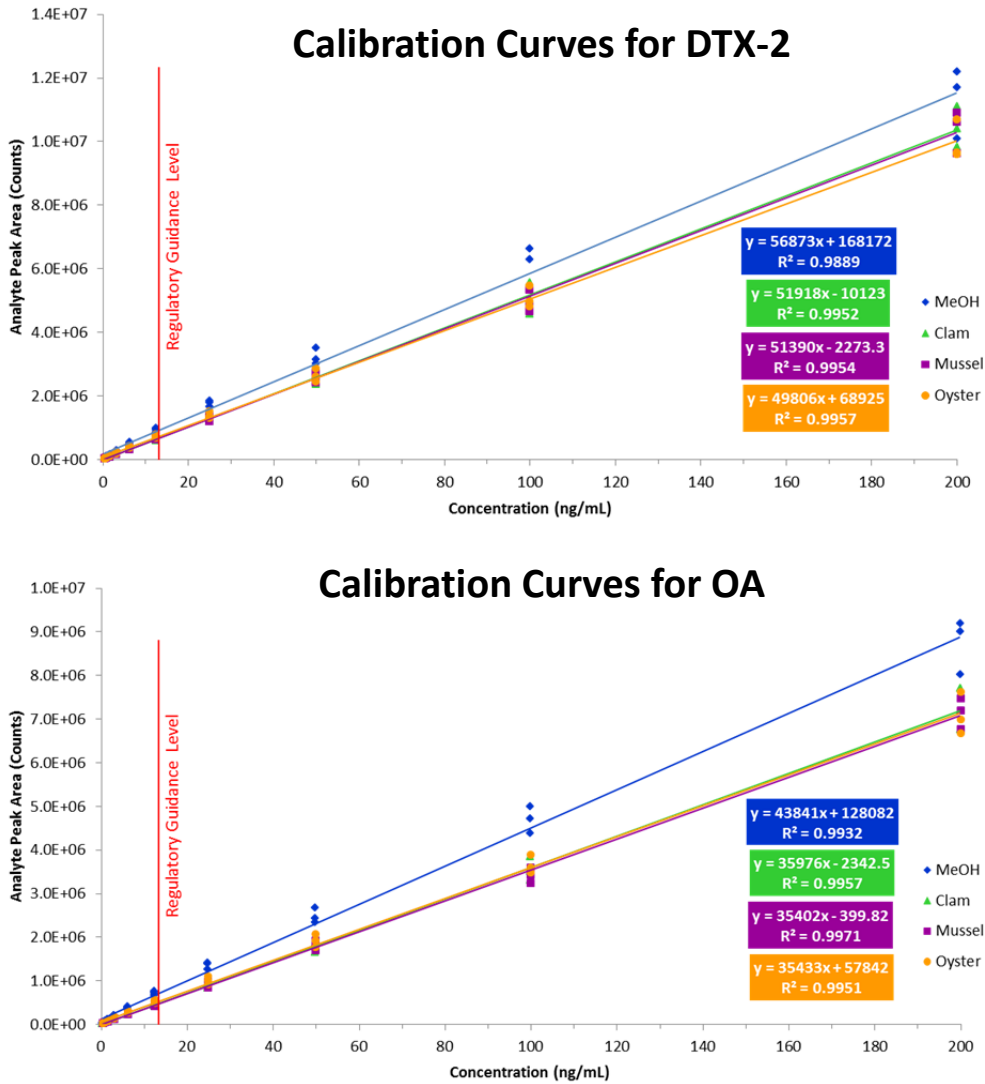


Figure 4. Calibration curves from neat standard solutions and spiked methanolic extracts of clam, mussel, and oyster analyzed under basic chromatographic conditions. A working range of 0.39–200 ng/mL (n=10) was used.

Pre-validation Limits of Detection (LODs) and Quantitation (LOQs):

LODs and LOQs are based on the standard deviation of the response and the slope.³ The equations for each are expressed as:

$$LOD = \frac{3.3\sigma}{S} \qquad LOQ = \frac{10\sigma}{S}$$

Where σ = the standard deviation of the response from five blank matrix samples
 S = the slope of the calibration curve

	LOD (ng/mL)			LOQ (ng/mL)		
	OA	DTX-2	DTX-1	OA	DTX-2	DTX-1
	Acidic / Basic	Acidic / Basic	Acidic / Basic	Acidic / Basic	Acidic / Basic	Acidic / Basic
Clam	0.055 / 0.040	0.010 / 0.003	0.032 / 0.043	0.166 / 0.120	0.031 / 0.009	0.096 / 0.129
Mussel	0.019 / 0.007	0.007 / 0.006	0.018 / 0.008	0.057 / 0.020	0.022 / 0.018	0.053 / 0.023
Oyster	0.017 / 0.015	0.008 / 0.011	0.018 / 0.016	0.050 / 0.046	0.025 / 0.034	0.054 / 0.049

Method Validation

Preparation of Stock Solutions for Validation:

Certified reference materials from the National Research Council Canada are supplied in sealed glass ampoules containing at least 0.5 mL of MeOH containing each toxin at a certified concentration that differs between toxins and lots. Thus, different volumes of each toxin standard, in 0.5 mL aliquots transferred using a 0.5 mL positive displacement Hamilton syringe, were transferred to a 20 mL glass scintillation vial and diluted with methanol to achieve stock standard solutions for each toxin at a concentration of 2 µg/mL. Stock solutions were stored at -20 °C.

Table 4. Toxin Stock Solution Preparation for Clam Validation Study

Certified Reference Material	Certified Concentration (µg/mL)	Lot #	Volume (mL)	Solvent (ml)	Total Volume	Final Concentration (µg/mL)
CRM-DTX1	15.1 ± 1.1	20071024	2	13.10	15.10	2
CRM-DTX2	7.8 ± 0.4	20071121	4	11.60	15.60	
CRM-OA-c	13.7 ± 0.6	20070328	2.5	14.625	17.125	

Validation Criteria:

Accuracy / Trueness, Measurement Uncertainty, Precision [Repeatability and Reproducibility], Recovery, Specificity, Working and Linear Ranges, Limit of Detection, Limit of Quantitation / Sensitivity, Ruggedness, Matrix Effects and Comparability (if intended as a substitute for an established method accepted by the NSSP).

Accuracy/Trueness

Each shellfish sample used for this study was screened for DSTs using the described LC-MS/MS method to assure that each matrix did not contain any naturally accumulated DSP toxins. Once these samples were established to be free of contamination, twenty sample homogenates (five each from matrix sources A through D from Table 1) were spiked at five concentrations ranging from 8 µg/100 g to 32 µg/100 g. Concentrations for spiking were selected to cover one half to two times the regulatory limit for DSTs in shellfish which is 16 µg of toxin per 100 g of shellfish tissue. Detailed procedures are described below.

1. For each sample, ten-twelve animals were rinsed, shucked, drained, and homogenized in a commercial food processor.
2. 2.0 ± 0.05 g of each homogenate was weighted into a 50 mL disposable centrifuge tube using a stainless steel laboratory spatula.
3. 9.0 mL of 100% methanol was added to the centrifuge tube

4. Individual DSP toxin standards were added into the tube, using positive displacement pipettes, so that the final concentration of each toxin was 8, 12, 16, 24, or 32 $\mu\text{g}/100\text{ g}$
5. Each sample was mixed for 3 min using a vortex mixer
6. Samples were centrifuged at 2000 *g* for 10 min at 20 °C
7. The supernatant was transferred to a 20 mL glass scintillation vial.
8. The residual tissue pellet was re-extracted with 9.0 mL of methanol and homogenized using a Polytron homogenizer, followed by vortexing for 3 min.
9. Samples were centrifuged at 2000 *g* for 10 min at 20 °C
10. The supernatant was transferred to a clean 25 mL graduated cylinder, combined with the first extract, and the total extract volume was adjusted to 20 mL by adding 100% methanol. The adjusted extract was then transferred back into the original 20 mL glass scintillation vial.
11. A 2 mL aliquot of the 20 mL extract was transferred to a 16 × 100 mm glass tube using a 1 mL positive displacement Hamilton syringe and 250 μL of 2.5 M NaOH was added. Each tube was sealed with a phenolic PTFE lined screw cap and vortexed to mix for 30 seconds. Tube weights were recorded, then placed in a 76 ± 2 °C water bath for 40 minutes.
12. Sample tubes were dried, allowed to cool to room temperature for 5–10 minutes, and re-weighed to assess any evaporative sample loss. No samples required volume adjustment during these experiments. Samples were then neutralized with 2.5 M HCL, followed by vortex mixing for 30 seconds.
13. Approx. 5 mL of hexane was added to each 2.5 mL hydrolyzed methanolic extract and vortexed for 30 seconds to mix. Samples were partitioned by centrifuging at 2,000 *g* for 10 min at 20 °C. Using a disposable glass Pasteur pipette, the upper hexane layer was removed and discarded into an appropriate waste container, and approximately 1 mL of the remaining methanolic extract (bottom layer) was transferred into a 1 mL disposable syringe equipped with a 13 mm, 0.2 μm syringe tip filter using a clean disposable glass Pasteur pipette. Each filtrate was collected directly into an LC-MS certified glass vial for analysis.

Data for OA, DTX1, and DTX2 in clam are reported in tables 4, 5, and 6 respectively. These data were collected on four different days over the course of nine days.

Precision and Recovery

Matrix fortification and extraction were conducted as described above for Accuracy/Trueness. However, each of the ten samples (five samples each from sources A and B from Table 1) was fortified at concentrations of 8, 16, and 32 $\mu\text{g}/100\text{ g}$. Even number samples, comprised of clams from two source locations, were prepared on the same day and ran within 24 h. Odd number samples also consisting of clams from the two sources were prepared and analyzed nine days later than the even number samples. Data for OA, DTX1, and DTX2 are reported in tables 7, 8, and 9, respectively. Precision is reported as percent relative standard deviation (%RSD).

Specificity

All three DSTs were analyzed in the presence of the potentially co-occurring lipophilic toxins azaspiracids (NRC-CRM-AZA 1, NRC-CRM-AZA 2, NRC-CRM-AZA 3), pectenotoxin (NRC-CRM-PTX2) and yessotoxin (NRC-CRM -YTX). For each sample three aliquots of blank tissue matrix were used. One aliquot served as a control blank, one sample contained a ½ action level spike (8 $\mu\text{g}/100\text{ g}$) of each DSTs, and one

contained the same concentration of DSTs and also a moderate to high concentration of each potential interfering compound. Five technical replicates of each aliquot, excluding the negative control blank, were analyzed. The specificity index is reported for each DST.

Working and Linear Ranges, Limit of Detection, Limit of Quantitation / Sensitivity

Matrix fortification and extraction were conducted as described above for Accuracy/Trueness. However, each of ten replicate clam samples (five each from source locations A and B from Table 1) were fortified at five concentrations spanning 50-150% of the range of interest (4, 8, 16, and 32, 48 $\mu\text{g}/100\text{ g}$). Two replicate injections of each fortified sample were analyzed. The linear range for OA, DTX1, and DTX2 extracted from fortified clam is shown in figures 5, 6, and 7. These data were collected on four different days over the course of ten days.

Ruggedness

In addition to pre-validation ruggedness testing of acidic versus basic chromatography and neat versus matrix matched calibration curves (data presented earlier in this report), two additional factors were assessed: 1. Effect of the hexane washing step on accuracy/trueness, and 2. Effect of using a different mass spectrometer of the same make and model on method performance.

To assess the effect of the hexane washing step on accuracy/trueness, two sub-samples from each of 10 extracts from previously spiked samples, representing two different matrix sources (5 samples each from matrix sources A and B), each spiked at 5 different concentrations bracketing the regulatory guidance level (8, 12, 16, 24, and 32 $\mu\text{g}/100\text{ g}$), were hydrolyzed following the procedure above. For each sample, one hydrolyzed extract was put through the above described hexane washing step, while the other was filtered straight into an LC vial for analysis. This entire procedure was repeated on different days so that in total 20 samples were tested both with and without the hexane washing step. The data handling procedures outlined in the Marine Biotxin and Non-MPN Based Microbiological Methods SOP for Ruggedness were used for reporting these results.

To assess the effect of using a different mass spectrometer of the same make and model on method performance (i.e. method transferability), ten samples (spiked at either 8, 16, or 32 $\mu\text{g}/100\text{ g}$) were analyzed using two different AB Sciex QTrap 5500 detectors equipped with a Turbo V ionization source (same make and model but purchased several years apart). The data handling procedures outlined in the Marine Biotxin and Non-MPN Based Microbiological Methods SOP for Ruggedness were used for reporting these results.

Matrix Effects

See Pre-Validation Study Results above.

Comparability

Method comparability, typically accomplished through comparison to a reference method, was not directly possible due to the fact that no reference method has been established under the NSSP. LC-MS/MS is the established reference method for the measurement of lipophilic shellfish toxins in the EU. The method described here is a version of the EU reference method optimized specifically for DSP toxins. One of the biggest differences between the method described here and the EU reference method is that the EU reference method is designed to detect multiple groups of lipophilic toxins in addition to DSP such as pectenotoxins, azaspiracids, and yessotoxins. But the analysis of these additional toxin groups in a single chromatographic run requires switching between positive and negative ion modes and initial analysis without sample hydrolysis as hydrolysis destroys several of these toxin groups. Analysis of total DSP toxins (free plus esterified) requires a second sample injection after alkaline hydrolysis of a sub-sample of the shellfish extract. Pectenotoxins and yessotoxins are not required to be analyzed for under the NSSP as they have not been proven to cause human illness, and while azaspiracids are required to be monitored, an optimized LC-MS/MS method for azaspiracids would be performed in positive ion mode and without sample hydrolysis. Therefore we are treating the LC-MS/MS method for azaspiracids as a separate method, even though the same extract, mobile phase, and equipment can be used for the analysis of both toxins. Different labs in the EU do run different versions of the “reference method” including some analyzing with acidic chromatographic conditions and some using basic, but pre-validation studies performed here found that analysis under basic chromatographic conditions would require the use of matrix matched standards. Furthermore, the two regulatory laboratories in the U.S. currently running a version of the EU reference LC-MS/MS method for lipophilic shellfish toxins as best available science are both using acidic chromatographic conditions. Therefore, running a different version of the EU reference LC-MS/MS method, such as basic chromatography or analysis without hydrolysis did not seem relevant for the method comparability requirement.

Analysis of certified reference materials would be another way to assess method performance as these naturally contaminated materials are certified to contain a known amount of all three of the target compounds. Until recently the only source of CRMs for DSP toxins, NRC Canada, only certified these materials for free toxins (i.e. toxins present pre-hydrolysis not taking into account the potential presence of fatty acid acyl ester shellfish metabolites (DTX3), which are known to be present in naturally contaminated shellfish samples. But new materials produced by NRC, both a frozen shellfish homogenate and a lyophilized material, are now provided with informational concentrations for total toxins (free plus esterified). Five aliquots of the frozen CRM-DSP-Mus-c (at \$185.00 each) and one aliquot of the lyophilized NRC-FDMT1 (at \$1,175 each) were purchased from the NRC Canada Certified References Materials Program. The frozen CRM contains $4 \text{ g} \pm 0.5 \text{ g}$ of homogenized material therefore can only be reliably tested once using the required 2 g extraction method. The lyophilized material reportedly contains enough material for approximately 8 extractions. This material was extracted and tested 5 times, for a total of 10 CRM replicate tests (5 frozen and 5 lyophilized). Each extract was hydrolyzed and tested twice on separate days (for a total of 20 analyses) to assess the methods performance.

Although LC-MS/MS is the only reference method currently accepted in the EU for the analysis of lipophilic shellfish toxins, EU regulations do allow for the use of supplementary methods if they are shown to be equally protective (Commission Regulations (EC) No.853/2004 and No.15/2011). One such supplementary method that has been both single⁶ and multi-laboratory⁷ validated and is recognized as equally protective for DSP toxins in the EU is the OkaTest, produced by ZEU Inmunotec in Spain. The colorimetric protein phosphatase inhibition assay (PPIA), OkaTest, complies with the criteria stipulated by the European Reference Laboratory on Marine Toxins and Commission Regulation 15/2012 for determination of OA-group toxins in molluscs, according to the European Commission (DG-SANCO). But since this method specifically detects DSP toxins while the EU reference LC-MS/MS method detects a number of different lipophilic shellfish toxins, the OkaTest is considered a supplementary method for the detection of DSP and not a full alternative to LC-MS/MS for the lipophilic shellfish toxins by the EU. The OkaTest is now sold in the United States by Abraxis LLC as the Protein Phosphatase Inhibition Assay (PPIA) for DSP toxins. Since the PPIA OkaTest provides a composite toxicity score for all DSP toxins present, it was not possible to test all of the spiked samples generated during this validation directly by both methods as all samples in this study were spiked with all three of the target compounds. Furthermore, the linear range of the OkaTest is 6.3 – 35.4 µg OA eq./100 g. Only two of the spiking levels used in this study were within the linear range of the OkaTest, the 4 and 8 µg/100 g spiking levels (n=10 each; 20 samples total) from the linear range determination, which equated to 12 and 24 µg/100 g in total DSP toxins, closely bracketing the regulatory level of 16 µg OA eq./100 g. In addition, all shellfish homogenates (n = 10 each; 30 samples total) from the three species of shellfish used in the pre-validation matrix effect studies (clams, *Mercinaria mercinaria*; mussels, *Mytilus edulus*; and oysters, *Crassostrea virginica*), that had been previously tested and found to be <LOD for DSP toxins by LC-MS/MS were also tested using the OkaTest to show that they were also negative by this alternative method. Lastly, naturally contaminated shellfish from a variety of species and geographic locations including softshell clams (*Mya arenaria*) from New York (n=9) and containing both OA and DTX1 (range <LOD – 37.3 µg OA eq. /100 g), blue mussels (*Mytilus edulus*) from Washington (n=12) containing DTX1 only (<LOD – 52.5 µg/100 g), and eastern oysters (*Crassostrea virginica*) from Texas (n=11) containing OA only (1.6 – 56.3 µg/100g) were tested by both methods to assess comparability of the LC-MS/MS method with PPIA.

Results

To be provided to the LMRC with sufficient time to be reviewed prior to the 2017 ISSC meeting.

ISSC Method Application and Single Lab Validation Checklist For Acceptance of a Method for Use in the NSSP

The purpose of single laboratory validation in the National Shellfish Sanitation Program (NSSP) is to ensure that the analytical method under consideration for adoption by the NSSP is fit for its intended use in the Program. A Checklist has been developed which explores and articulates the need for the method in the NSSP; provides an itemized list of method documentation requirements; and, sets forth the performance characteristics to be tested as part of the overall process of single laboratory validation. For ease in application, the performance characteristics listed under validation criteria on the Checklist have been defined and accompany the Checklist as part of the process of single laboratory validation. Further a generic protocol has been developed that provides the basic framework for integrating the requirements for the single laboratory validation of all analytical methods intended for adoption by the NSSP. Methods submitted to the Interstate Shellfish Sanitation Conference (ISSC) Laboratory Methods Review (LMR) Committee for acceptance will require, at a minimum, six (6) months for review from the date of submission.

Name of the New Method	Liquid Chromatography Tandem Mass Spectrometry (LC-MS/MS) Method for the Determination of Diarrhetic Shellfish Poisoning (DSP) Toxins in Shellfish.	
Name of the Method Developer	Jonathan Deeds	
Developer Contact Information	Jonathan.deeds@fda.hhs.gov ; 240-402-1474 US FDA, 5001 Campus Drive, College Park, MD 20740	
Checklist	Y/N	Submitter Comments
A. Need for the New Method		
1. Clearly define the need for which the method has been developed.	Y	
2. What is the intended purpose of the method?	Y	
3. Is there an acknowledged need for this method in the NSSP?	Y	
4. What type of method? i.e. chemical, molecular, culture, etc.	Y	
B. Method Documentation		
1. Method documentation includes the following information:	Y	
Method Title	Y	
Method Scope	Y	
References	Y	
Principle	Y	
Any Proprietary Aspects	Y	
Equipment Required	Y	
Reagents Required	Y	
Sample Collection, Preservation and Storage Requirements	Y	
Safety Requirements	Y	
Clear and Easy to Follow Step-by-Step Procedure	Y	
Quality Control Steps Specific for this Method	Y	
C. Validation Criteria		
1. Accuracy / Trueness	Y	
2. Measurement Uncertainty	Y	
3. Precision Characteristics (repeatability and reproducibility)	Y	
4. Recovery	Y	
5. Specificity	Y	
6. Working and Linear Ranges	Y	
7. Limit of Detection	Y	

8. Limit of Quantitation / Sensitivity	Y	
9. Ruggedness	Y	
10. Matrix Effects	Y	

11. Comparability (if intended as a substitute for an established method accepted by the NSSP)	Y	
D. Other Information		
1. Cost of the Method	Y	
2. Special Technical Skills Required to Perform the Method	Y	
3. Special Equipment Required and Associated Cost	Y	
4. Abbreviations and Acronyms Defined	Y	
5. Details of Turn Around Times (time involved to complete the method)	Y	
6. Provide Brief Overview of the Quality Systems Used in the Lab	Y	
Submitters Signature		
Jonathan R. Deeds -S <small>Digitally signed by Jonathan R. Deeds -S DN: c=US, o=U.S. Government, ou=HHS, ou=FDA, ou=People, 0.9.2342.19200300.100.1.1=1300218767, cn=Jonathan R. Deeds -S Date: 2017.05.31 12:46:30 -04'00'</small>	Date:	5/31/2017
Submission of Validation Data and Draft Method to Committee		
Date:		
Reviewing Members		
Date:		
Accepted		
Date:		
Recommendations for Further Work		
Date:		
Comments:		

DEFINITIONS

1. **Accuracy/Trueness** - Closeness of agreement between a test result and the accepted reference value.
2. **Analyte/measurand** - The specific organism or chemical substance sought or determined in a sample.
3. **Blank** - Sample material containing no detectable level of the analyte or measurand of interest that is subjected to the analytical process and monitors contamination during analysis.
4. **Comparability** - The acceptability of a new or modified method as a substitute for an established method in the NSSP. Comparability must be demonstrated for each substrate or tissue type by season and geographic area if applicable.
5. **Fit for purpose** - The analytical method is appropriate to the purpose for which the results are likely to be used.
6. **HORRAT value** - HORRAT values give a measure of the acceptability of the precision characteristics of a method.⁴
7. **Limit of Detection** - the minimum concentration at which the analyte or measurand can be identified. Limit of detection is matrix and analyte/measurand dependent.⁴
8. **Limit of Quantitation/Sensitivity** - the minimum concentration of the analyte or measurand that can be quantified with an acceptable level of precision and accuracy under the conditions of the test.
9. **Linear Range** - the range within the working range where the results are proportional to the concentration of the analyte or measurand present in the sample.
10. **Measurement Uncertainty** - A single parameter (usually a standard deviation or confidence interval) expressing the possible range of values around the measured result within which the true value is expected to be with a stated degree of probability. It takes into account all recognized effects operating on the result including: overall precision of the complete method, the method and laboratory bias and matrix effects.
11. **Matrix** - The component or substrate of a test sample.
12. **Method Validation** - The process of verifying that a method is fit for purpose.¹
13. **Precision** - the closeness of agreement between independent test results obtained under stipulated conditions.^{1, 2} There are two components of precision:
 - a. **Repeatability** - the measure of agreement of replicate tests carried out on the same sample in the same laboratory by the same analyst within short intervals of time.
 - b. **Reproducibility** - the measure of agreement between tests carried out in different laboratories. In single laboratory validation studies reproducibility is the closeness of agreement between results obtained with the same method on replicate analytical portions with different analysts or with the same analyst on different days.
14. **Quality System** - The laboratory's quality system is the process by which the laboratory conducts its activities so as to provide data of known and documented quality with which to demonstrate regulatory compliance and for other decision-making purposes. This system includes a process by which appropriate analytical methods are selected, their capability is evaluated, and their performance is documented. The quality system shall be documented in the laboratory's quality manual.
15. **Recovery** - The fraction or percentage of an analyte or measurand recovered following sample analysis.
16. **Ruggedness** - the ability of a particular method to withstand relatively minor changes in analytical technique, reagents, or environmental factors likely to arise in different test environments.⁴
17. **Specificity** - the ability of a method to measure only what it is intended to measure.¹
18. **Working Range** - the range of analyte or measurand concentration over which the method is applied.

REFERENCES:

1. Eurachem Guide, 1998. The Fitness for Purpose of Analytical Methods. A Laboratory Guide to Method Validation and Related Topics. LGC Ltd. Teddington, Middlesex, United Kingdom.
2. IUPAC Technical Report, 2002. Harmonized Guidelines for Single-Laboratory Validation of Methods of Analysis, Pure Appl. Chem., Vol. 74, (5): 835-855.
3. Joint FAO/IAEA Expert Consultation, 1999. Guidelines for Single-Laboratory Validation of Analytical Methods for Trace-Level Concentrations of Organic Chemicals.
4. MAF Food Assurance Authority, 2002. A Guide for the Validation and Approval of New Marine Biotxin Test Methods. Wellington, New Zealand.
5. National Environmental Laboratory Accreditation. , 2003. Standards. June 5.
6. EPA. 2004. EPA Microbiological Alternate Procedure Test Procedure (ATP) Protocol for Drinking Water, Ambient Water, and Wastewater Monitoring Methods: Guidance. U.S. Environmental Protection Agency (EPA), Office of Water Engineering and Analysis Division, 1200 Pennsylvania Avenue, NW, (4303T), Washington, DC 20460. April.

ISSC Method Application and Single Lab Validation Checklist For Acceptance of a Method for Use in the NSSP

The purpose of single laboratory validation in the National Shellfish Sanitation Program (NSSP) is to ensure that the analytical method under consideration for adoption by the NSSP is fit for its intended use in the Program. A Checklist has been developed which explores and articulates the need for the method in the NSSP; provides an itemized list of method documentation requirements; and, sets forth the performance characteristics to be tested as part of the overall process of single laboratory validation. For ease in application, the performance characteristics listed under validation criteria on the Checklist have been defined and accompany the Checklist as part of the process of single laboratory validation. Further a generic protocol has been developed that provides the basic framework for integrating the requirements for the single laboratory validation of all analytical methods intended for adoption by the NSSP. Methods submitted to the Interstate Shellfish Sanitation Conference (ISSC) Laboratory Methods Review (LMR) Committee for acceptance will require, at a minimum, six (6) months for review from the date of submission.

Name of the New Method		
Matrix Expansion for the Receptor Binding Assay (RBA) for Paralytic Shellfish Poisoning (PSP) Toxicity Determination to Allow Use with Geoduck		
Name of the Method Developer		
Michael Jamros, Chris Whitehead		
Developer Contact Information		
Sitka Tribe of Alaska, 456 Katlian St, Sitka, AK 99835		
907-747-7356 phone		
907-747-4915 fax		
michael.jamros@sitkatriben-sns.gov		
chris.whitehead@sitkatriben-sns.gov		
Checklist		
A. Need for the New Method		
1. Clearly define the need for which the method has been developed.		
<p>Paralytic shellfish poisoning (PSP) is a food born illness caused by ingestion of contaminated shellfish. The paralytic shellfish toxin, saxitoxin (STX), and its analogs are potent neurotoxins responsible for PSP. Marine dinoflagellates and freshwater cyanobacteria produce STX. The STX can accumulate in filter-feeding bivalve mollusks to levels that are toxic to humans. Symptoms of PSP include: tingling and numbness of the perioral area and extremities, drowsiness, incoherence, loss of motor control, and following high dose consumption, respiratory paralysis.</p> <p>In 1965 the mouse bioassay (MBA) was adopted as an official AOAC method for STX determination. The MBA has served as the primary method available for PSP testing for the last five decades. Both North American and European regulatory agencies have expressed the desire to transition to a more humane PSP testing method that does not require the use of live animals and is not subject to the matrix effects documented for the MBA (Turner 2012). Recently, the NSSP approved a post-column oxidation liquid chromatographic (PCOX HPLC) method and a receptor binding assay (RBA) as alternatives to the MBA. The PCOX HPLC method is approved for full use; whereas, the RBA is approved for limited use (the RBA is only approved for shellfish matrices evaluated in the single lab and multi-lab validation studies, which does not include geoduck (<i>Panopea</i>). Both the PCOX and RBA are sensitive quantitative assays for STX detection, and they do not require the use of live animals. The PCOX HPLC requires skilled personnel and offers low throughput in comparison to the RBA.</p>		
2. What is the intended purpose of the method?		
<p>The RBA is approved for regulatory testing of mussels as an alternative to the MBA and is approved for limited use as a screening tool for clams and scallops, but is not yet approved for use with geoduck (<i>Panopea</i>) due to a lack of data. Geoduck are a major commercial product that requires PSP testing. This proposal requests consideration for the NSSP RBA approval to be expanded to include geoduck. The proposal provides data from a single laboratory validation (SLV) of the RBA for geoduck testing as support for this request.</p>		

<p>This method is intended for use as an NSSP Approved Limited Use Method for screening for PSP toxicity in shellfish, specifically geoducks. The RBA serves as an alternative to the MBA in these applications, offering a measure of integrated toxicity with high throughput and the elimination of live animal testing (Van Dolah 2013). This application is for the addition of geoduck to the list of matrices approved for use with the RBA.</p>
<p>3. Is there an acknowledged need for this method in the NSSP?</p> <p>There is an acknowledged need for this method extension in the NSSP. A significant portion of the Washington and Alaska state shellfish industries are comprised of the harvest of geoduck. Approval of the RBA for use with geoduck would provide an alternative to (1) the MBA, which uses live animals, and (2) the PCOX HPLC method, which requires costly equipment and skilled personnel and offers low throughput.</p> <p>Acceptance of the RBA as an NSSP Approved Method for Marine Biotoxin Testing for PSP toxicity determination in geoduck would provide monitoring and management programs with an additional tool that can be used for monitoring toxin levels and making regulatory decisions. Not only does the RBA eliminate the need for live animals for PSP testing, it is also more sensitive than the MBA.</p>
<p>4. What type of method? i.e. chemical, molecular, culture, etc.</p> <p>Molecular. The RBA is a functional assay, whereby toxins present in the standard/sample bind to sodium channel preparations in the assay. Radiolabeled toxins (3H-STX) compete with toxins present in the standard or sample for sodium channel binding sites in a microplate format. Thus a decrease in signal from radiolabeled toxins represents an increase in standard/sample toxicity. This competitive RBA allows for quantitation that directly relates to the composite toxicity of the sample.</p>
<p>B. Method Documentation</p>
<p>1. Method documentation includes the following information:</p>
<p>Method Title</p> <p>Matrix Expansion for the Receptor Binding Assay (RBA) for Paralytic Shellfish Poisoning (PSP) Toxicity Determination to Allow Use with Geoduck</p>
<p>Method Scope</p> <p>This submission presents the ‘Matrix Expansion for the Receptor Binding Assay (RBA) for Paralytic Shellfish Poisoning (PSP) Toxicity Determination to Allow Use with Geoduck’ for consideration as an NSSP Approved Method for Marine Biotoxin Testing for PSP in Geoduck.</p> <p>The RBA offers a high-throughput, sensitive, and quantitative alternative to the mouse bioassay (MBA), which has been the long-standing reference method for PSP toxicity. Further, the RBA eliminates the use of live animals for detection of these toxins. While the RBA still uses receptors prepared from animals, the number of animals required for analysis is significantly reduced. Using native receptors as the analytical recognition elements for the assay allows for a composite measure of overall toxicity, as opposed to toxin concentrations measured by liquid chromatographic methods that require conversion factors of equivalent toxicity to calculate the overall toxicity.</p> <p>The RBA has undergone AOAC single- and multi-laboratory validation and is designated through AOAC as an Official Method of Analysis (OMA 2011.27). The RBA is currently an NSSP Approved Method for Marine Biotoxin Testing for PSP in mussels as well as a NSSP approved for Limited Use Method for clams and scallops for the purpose of screening and precautionary closure for PSP (ISSC 2015 Summary of Actions Proposal 13-114). Here we provided results from a single laboratory validation study for use of RBA with the matrix geoduck viscera for submission for the RBA to be considered for approval as an NSSP Approved Method for Marine Biotoxin Testing for PSP.</p>
<p>References</p> <p>Van Dolah 2013. ISSC application: Receptor Binding Assay (RBA) for Paralytic Shellfish Poisoning (PSP) Toxicity Determination.</p> <p>Van Dolah et al. 2012. Determination of paralytic shellfish toxins in shellfish by receptor binding assay: collaborative study. J AOAC Int. May-Jun;95(3):795-812.</p>

<p>Van Dolah et al. 2009. Single-laboratory validation of the microplate receptor binding assay for paralytic shellfish toxins in shellfish. J AOAC Int. Nov-Dec;92(6):1705-13.</p> <p>Ruberu et al. 2012. Evaluation of variability and quality control procedures for a receptor-binding assay for paralytic shellfish poisoning toxins. Food AdditContam Part A Chem Anal Control Expo Risk Assess.29(11):1770-9.</p> <p>Turner et al. 2012. Investigations into matrix components affecting the performance of the official bioassay reference method for quantitation of paralytic shellfish poisoning toxins in oysters. Toxicon: official journal of the International Society on Toxicology 59, 215-230.</p> <p>OMA 2011.27. AOAC Official Method 2011.27 Paralytic shellfish toxins (PSTs) in shellfish, receptor binding assay. In Official Methods of Analysis of AOAC International. http://www.eoma.aoc.org.</p>
<p>Principle</p> <p>The RBA is a competition-based assay that employs radiolabeled Saxitoxin (3H-STX) to compete with PSP toxins present in standards/samples for binding sites on natural receptors in the assay. Following incubation with the receptors, unbound 3H-STX is removed and the remaining labeled toxin is measured with a scintillation counter. The amount of remaining 3H-STX is inversely proportional to standard/sample toxicity.</p>
<p>Any Proprietary Aspects</p> <p>None</p>
<p>Equipment Required</p> <p>The following list identifies the equipment and supplies needed for conducting the RBA.</p> <p>For the assay:</p> <ul style="list-style-type: none"> (a) Scintillation counter (traditional or microplate) (b) An 8-channel pipettor (5-200 ul variable volume and disposable tips) (c) Micropipettors (1-1000 ul variable volumes and disposable tips) (d) 96-well microtitre filter plate (1 µm pore size type GF/B glass fiber filter/0.65 um pore size Durapore support membrane (Millipore, Bedford, MA; Cat. No. MSFB N6B 50) (e) MultiScreen vacuum manifold (Millipore; Cat. No. NSVMHTS00) (f) Vacuum pump (g) Centrifuge tubes (15 and 50 ml, conical, plastic) (h) Mini dilution tubes in 96-tube array (i) Reagent reservoirs (j) Ice bucket and ice (k) Vortex mixer (l) Sealing tape (Millipore; Cat. No. MATA HCL00) (m) Volumetric flask or graduated beaker (1 L) (n) -80 °C freezer (o) Refrigerator <p>For sample extraction:</p> <ul style="list-style-type: none"> (p) Blender or homogenizer for sample homogenization (q) Pipets (r) Centrifuge tubes (15 ml, conical, plastic) (s) pH meter or pH paper (t) Hot plate or water bath (u) Graduated centrifuge tubes (15 ml) (v) Centrifuge and rotor for 15 ml tubes
<p>Reagents Required</p> <p>For the assay:</p> <ul style="list-style-type: none"> a) STX diHCl standards (NIST RM 8642; available through the National Institute of Standards and Technology; www.nist.gov) [This is the same standard used for the MBA] or (CRM-STX; National Research Council of Canada;

<p>www.nrc-cnrc.gc.ca/eng/solutions/advisory/crm/list_product.html#B-PSP)</p> <p>(b) 3H-STX (0.1 mCi per ml, ≥10 Ci per mmol; available through American Radiolabeled Chemicals, St. Louis, MO [or equivalent])</p> <p>(c) 3-Morpholinopropanesulfonic acid (MOPS; Sigma; St. Louis, MO; Cat. No. M3183-500G [or equivalent])</p> <p>(d) Choline chloride (Sigma; Cat. No. C7527-500G [or equivalent])</p> <p>(e) Ultima Gold liquid scintillation cocktail (PerkinElmer Inc.; Waltham, MA; Cat. No. 6013321 [or equivalent])</p> <p>For the sample extraction:</p> <p>(f) Hydrochloric acid (HCl; 1.0 and 0.1 M)</p> <p>(g) Sodium hydroxide (0.1 M)</p> <p>(h) Water (distilled or deionized [18 µΩ])</p>
<p>Sample Collection, Preservation and Storage Requirements</p> <p>Samples should be kept cool until meat is removed from shell, meat should be removed from shell within 48 hours of collection and either frozen or extracted.</p>
<p>Safety Requirements</p> <p>General safety requirements (e.g., personal protective equipment including gloves, safety glasses, and laboratory coat) for working with toxins, biological reagents, and radioactive material must be followed. Users must be trained in and follow all in-house safety procedures for working with toxins and radiolabeled materials. Even though low levels of radiation are used for this assay, users must follow all local, state and federal laws and procedures regarding the receipt, use, and disposal of isotopes.</p>
<p>Clear and Easy to Follow Step-by-Step Procedure</p> <p>Please see the detailed protocol AOAC OMA 2011.27 (Appendix 1)</p>
<p>Quality Control Steps Specific for this Method</p> <p>Only data falling within the linear part of the curve (0.2-0.7 B/B₀) is used for quantitation. Binding curve data shown here is from 14 RBA plates run on separate days. All analysis was performed using GraphPad Prism version 7.02.</p> <p>The following parameters are required for quality control and acceptance of RBA results and were met by all assays included in this study:</p> <ul style="list-style-type: none"> (a) Slope must be between -0.8 and -1.2 (theoretical slope is -1). In this study, the average slope was -0.98 +/- 0.08. (b) IC₅₀ (inhibitory concentration at which CPM is 50% maximum) is in the acceptable range (2.0 nM ± 30%), between 1.4 and 2.6 nM. In this study, the average IC₅₀ was 1.7 nM +/- 0.1 nM. (c) A QC sample (1.8 x 10⁻⁸ M STX concentration, 3 nM STX in-well concentration) should be within 30% (2.1 nM to 3.9 nM in-well concentration). In this study, the measured QC had an average value of 3.1 nM +/- 0.4 nM. (d) The RSDs of triplicate counts per minute must be less than 30%. All standards, QC samples, and geoduck samples in this study met these criteria.
<p>C. Validation Criteria</p>
<p>1. Accuracy / Trueness</p> <p>Accuracy was evaluated based on recovery of known amounts of saxitoxin added as a QC check sample. A QC check sample is included in every receptor binding assay. Recovery of the QC check sample (3nM in-well solution) was 105% +/- 13% (Table 1).</p>

Table 1: Calibration curve and QC check parameters in receptor binding assays

RBA ID	Slope	R ²	IC ₅₀ (nM)	IC ₇₀ (nM)	LOQ (ug STX eq/ 100g tissue)	QC (nM)
17-001	-0.86	0.99	1.6	0.59	2.6	2.9
17-002	-0.88	0.99	1.8	0.68	3.0	2.8
17-003	-0.94	0.96	1.6	0.65	2.9	2.5
17-004	-0.99	0.96	1.7	0.71	3.2	2.6
17-005	-0.92	0.98	1.5	0.60	2.7	3.1
17-006	-0.98	0.98	1.8	0.78	3.5	3.1
17-009	-0.95	0.94	1.5	0.62	2.8	3.6
17-010	-1.00	0.96	1.5	0.66	2.9	3.0
17-011	-1.15	0.96	1.9	0.92	4.1	3.7
17-012	-1.08	0.97	1.7	0.77	3.4	3.3
17-013	-1.04	0.97	1.8	0.81	3.6	3.1
17-014	-0.99	0.95	1.7	0.70	3.1	3.1
17-015	-0.95	0.99	1.5	0.62	2.8	3.7
17-016	-1.04	0.96	1.8	0.77	3.4	3.4
Average	-0.98	0.97	1.7	0.71	3.2	3.1
+/-	0.08	0.02	0.1	0.09	0.4	0.4

2. Measurement Uncertainty

3. Precision Characteristics (repeatability and reproducibility)

Repeatability was determined by analyzing each sample in three assays performed on independent days. The average RSD was 14.6%, with a range of 5.4% to 25.6% (Table 2). These results are consistent with the mean RSD of 17.7% (Van Dolah 2009), used to demonstrate repeatability in ISSC 2015 Proposal 13-114.

Table 2: Receptor binding assay results, summary statistics, and comparison to MBA results

Sample ID	RBA (ug/100g)			RBA mean (ug/100g)	MBA (ug/100g)	% MBA value	SD	RSD (%)
1823	29	32	38	33	42	79	5	13.9
2095	22	37	34	31	45	69	8	25.6
1594	45	74	58	59	58	102	15	24.6
2094	51	56	48	52	59	88	4	7.8
1607	60	43	47	52	67	78	12	23.3
1865	88	111	86	95	75	127	14	14.6
1933	88	85	74	82	88	93	7	9
1830	121	108	83	104	116	90	19	18.6
2315	93	97	82	91	128	71	8	8.6
2420	103	98	111	104	129	81	7	6.3
2071	129	141	163	144	140	103	17	11.9
2072	169	152	158	160	142	113	9	5.4
2138	406	344	332	361	447	81	40	11
1595	25	31	19	25	<38	-	6	24
1674	3	9	6	6	NTD	-	3	50*
Average						90	12	14.6

*RSD value omitted due to value below LOQ

4. Recovery

The average recovery of the QC check sample (3 nM in-well solution) was 105% +/- 13%.

5. Specificity

The RBA is specific to toxins that bind to site 1 of voltage-gated sodium channels. This includes all PSP congeners, whereby binding affinity is proportional to potency. Tetrodotoxin also binds to site 1 of the sodium channels, yet the typical combinations of sources, vectors, and geographical regions of tetrodotoxin and the saxitoxins differ.

6. Working and Linear Ranges

The dynamic range of the RBA is 1.2-10.0 nM in-well concentration (Van Dolah 2012). When necessary, samples must be diluted prior to analysis so that they are within the dynamic range of the RBA. Sigmoidal dose response with variable slope analysis is used to generate a binding curve from standard STX concentrations evaluated on each plate.

7. Limit of Detection

See Table 3 in the next section for a description of the limit of detection (LOD) for this method

8. Limit of Quantitation / Sensitivity

The limit of quantitation (LOQ) was determined from the average IC70 of all assays ran in the study, which was 0.71 nM +/- 0.09 nM. Using an adaptation of Eurachem Guide definitions for limit of detection (LOD) and LOQ by Van Dolah et. al. (2012), where B/B0 = 0.7 (average IC70 value) is used as the cutoff for quantitation, we obtain the below values for LOD and LOQ (Table 1). The numbers are for a sample diluted 1/10 (the established minimum dilution to avoid matrix effects) and extraction according to the AOAC protocol.

Table 3: LOD and LOQ for RBA matrix expansion of geoduck SLV

	Equation	SLV Results
LOD	$IC_{70} + 3 \times SD$	4.4 ug STX eq/100 g
LOQ	$IC_{70} + 10 \times SD$	7.2 ug STX eq/100 g

9. Ruggedness

Previous work has been done to identify critical steps to ensure accuracy and ruggedness (Ruberu et al. 2012, Van Dolah et al. 2012, VanDolah et al. 2009). It was deemed important to clarify the shellfish extracts by centrifugation prior to performing the assay, particularly if the sample was refrigerated or frozen. The rat brain preparations should be vortexed frequently to ensure the synaptosomes are in suspension, and the buffer should be ice cold to ensure that toxins are not released from the receptor. Assay plate filtration should be at a rate of 2-5 seconds for well clearance. Lastly, a minimum of 30 minutes should be allowed before reading the plates after scintillation liquid is added such that scintillant can penetrate the filters (Van Dolah 2013).

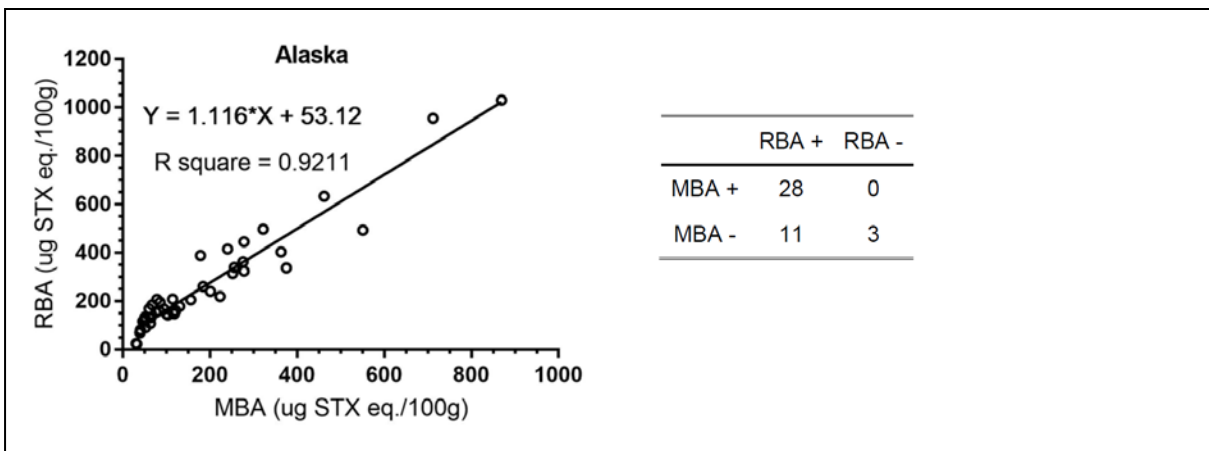
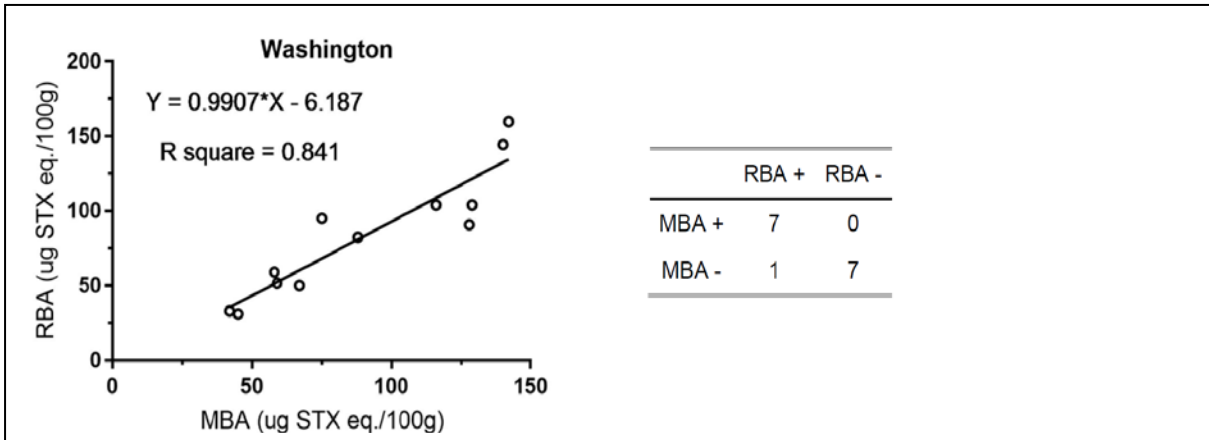
10. Matrix Effects

No matrix effects were reported. Minimum dilutions of shellfish extracts were 10-fold and were found to be sufficient to eliminate matrix effects. (Van Dolah 2013)

11. Comparability (if intended as a substitute for an established method accepted by the NSSP)

Comparability to MBA

A comparison of STX concentration assayed in naturally contaminated samples by the MBA and the RBA was performed using linear regression analysis (GraphPad Prism, version 7.02). MBA results for samples from Washington were analyzed by the Washington Department of Health Shellfish Biotoxins & Water Bacteriology Laboratories and samples from Alaska were analyzed by the Alaska Department of Environmental Conservation Environmental Health Lab. All RBA results are from analysis by the Sitka Tribe of Alaska Environmental Research Laboratory. 57 total samples were compared, with the RBA yielding no false negatives relative to the regulatory limit of 80 ug/100g. Overall there were 12 false positives relative to the MBA.



Comparability to Previous RBA Validation Work

Previous work by (Van Dolah et al. 2012, Van Dolah et al. 2009) was submitted to the ISSC as ISSC 2015 Proposal 13-114, resulting in approval of the RBA as a NSSP Approved Method for PSP in mussels and as a NSSP Approved Limited Use Method for clams and scallops for the purpose of screening and precautionary closure for PSP. The results from this SLV for matrix expansion of RBA for geoduck matrix is consistent with the data from the previous validation studies.

A comparison of this SLV to previous validation work for the RBA demonstrates the ability of the RBA to withstand minor changes in analytical technique, reagents, and environmental factors (Table 4).

Table 4: Comparison of SLV results to previous RBA validation studies						
	Accuracy (recovery of QC)	Repeatability (Average RSD)	Linear Range (slope, R ²)	IC ₅₀ (nM)	LOQ (mean IC ₇₀ - nM)	Comparison to MBA (R ² from linear regression analysis)
STA Geoduck Van Dolah et. al. 2009 - SLV	104.5%	14.6%	-0.98, 0.97	1.7 +/- 0.1	0.7	0.84, 0.92
Van Dolah et. al. 2012 - MLV	99.3%	17.1%	-0.98, 0.97	2.3 +/- 0.3	1.1	0.98, 0.88
	106.9%	17.1%	-1.03, ND*	1.9 +/- 0.5	0.8	0.84
*No data available						
D. Other Information						
1. Cost of the Method						
The estimated cost per 96-well plate assay is ~\$95.00. Including standards and samples with triplicate measurements (as well as three dilutions per sample [ranging from 3.5-600 µg STX eq 100 g ⁻¹] to ensure the unknown samples fall within linear range of assay), the cost per sample for quantitation would be ~\$13.60. If running multiple plates or in screening mode, sample costs would be reduced. (Van Dolah 2013)						
2. Special Technical Skills Required to Perform the Method						
General laboratory training is necessary (this would include being able to prepare reagent solutions, pipetting, centrifugation, and simple calculations). Additional training for working with low levels of radioactive material is required.						
3. Special Equipment Required and Associated Cost						
A microplate scintillation counter is needed and the cost is ~\$50-120K for a new counter, depending on the brand and number of simultaneous detectors. However, used instruments can be purchased for ~\$15K.						
4. Abbreviations and Acronyms Defined						
3H-STX Tritiated saxitoxin						
AOAC, Association of Analytical Communities						
ARC, American Radiolabeled Chemicals						
B, Bound CPM						
Bo, Maximum bound CPM						
CFSAN, Center for Food Safety & Applied Nutrition						
CPM, Counts per minute						
diHCl, Dihydrochloride						
Eq, Equivalent						
HCl, Hydrochloric acid						
IC50, Inhibitory concentration at which CPMs are at 50% of maximum						
LC-FD, Liquid chromatography with fluorescence detection						
LOD, Limit of detection						
LOQ, Limit of quantitation						
MBA, Mouse bioassay						
MOPS, 3-Morpholinopropanesulfonic acid						
NaOH, Sodium hydroxide						
NIST, National Institute of Standards and Technology						
NSSP, National Shellfish Sanitation Program						
OMA, Official method of analysis						
PCOX, Post-column oxidation liquid chromatography with fluorescence detection						
Pre-COX, Pre-column oxidation liquid chromatography with fluorescence detection						
PSP, Paralytic shellfish poisoning						
PSTs, Paralytic shellfish toxins						
QC, Quality control						

<p>QS, Quality System RBA, Receptor binding assay RSD, Relative standard deviation SLV, Single laboratory validation STX, Saxitoxin</p>
<p>5. Details of Turn Around Times (time involved to complete the method) Microplate scintillation counting provides the ability to test multiple samples simultaneously with a turn around time for data in approximately 3 hours. Up to six plates per analyst are possible in one day, yielding a throughput of 42 samples per day. If the assay is run in screening mode where only a single dilution (1/10) is run, then through-puts of >120 samples per day can be achieved.</p>
<p>6. Provide Brief Overview of the Quality Systems Used in the Lab The Center for Food Safety and Applied Nutrition (CFSAN) Quality System (QS) provides guidance to (1) design and develop processes, products, and services related to CFSAN’s mission, the FDA’s regulatory mission, and critical management and administrative support services, and (2) continually improve and strengthen product and service quality. The Laboratory Quality Assurance program serves as CFSAN’s logical application of QS to Center laboratories and lab-based activities. The third edition (October 2009) of the Laboratory Quality Manual was followed. Standard reference materials for saxitoxin are obtained through the National Institute of Standards and Technology (NIST) and are accompanied by a Report of Investigation. The standard reference saxitoxin used in the RBA is the same as that employed with the MBA. The 3H-STX is obtained through American Radiolabeled Chemicals, Inc., and is accompanied by a Technical Data Sheet with lot specifications.</p>

Appendix 1

AOAC Official Method 2011.27 Paralytic Shellfish Toxins (PSTs) in Shellfish Receptor Binding Assay First Action 2011

[Applicable to the determination of paralytic shellfish toxins (PSTs), as μg STX diHCl equiv./kg, in shellfish (mussels, clams, scallops) at levels $>149 \mu\text{g}$ STX diHCl equiv./kg, with a limit of detection (LOD) of 45 STX diHCl equiv./kg shellfish and a limit of quantification (LOQ) of 126 μg STX diHCl equiv./kg shellfish.]

Caution: Wear disposable gloves and protective laboratory coat while performing the assay. PSTs are neurotoxins that are harmful if ingested. The assay uses a tritium labeled tracer, [^3H] STX, at low concentration. All laboratories performing the assay must have approved radiation laboratory space and must follow procedures prescribed by their nuclear regulatory agency for receipt, use, and disposal of isotopes.

See Tables 2011.27A–E for the results of the interlaboratory study supporting acceptance of the method.

A. Principle

Test portions of shellfish homogenates are extracted using the AOAC mouse bioassay extraction protocol (959.08), modified by scale. The PST receptor assay is a competitive binding assay in which [^3H] STX competes with unlabeled STX in standards or mixtures of PST in samples for a finite number of available receptor sites (site 1 on the voltage gated sodium channel) in a rat brain membrane preparation. Following establishment of binding equilibrium at 4°C, unbound [^3H] STX is removed by filtration and bound [^3H] STX is quantified by liquid scintillation counting. A standard curve is generated using increasing concentrations of STX standard from 10^{-10} to 10^{-6} M STX, which results in a reduction in bound [^3H] STX that is directly proportional to the amount of unlabeled toxin present. The concentration of toxin in samples is determined in reference to the standard curve. Incubation is carried out in a microplate format to minimize sample handling and the amount of radioactivity used. Bound [^3H] STX (as counts per minute; CPM) can be determined either by conventional or by microplate scintillation counting. Both methods are included in this protocol.

B. Apparatus and Supplies

- (a) *Traditional or microplate scintillation counter.*
- (b) *Micropipettors.*—1–1000 μL variable volumes and disposable tips.
- (c) *Eight channel pipettor.*—5–200 μL variable volume and disposable tips.
- (d) *96-Well microtiter filter plate.*—With 1.0 μm pore size type GF/B glass fiber filter/0.65 μm pore size Durapore support membrane (Millipore, Bedford, MA, USA; Cat. No. MSFB N6B 50).
- (e) *MultiScreen vacuum manifold.*—Millipore; Cat. No. NSVMHTS00.
- (f) *Vacuum pump.*
- (g) *Centrifuge tubes.*—15 and 50 mL, conical, plastic.
- (h) *Mini dilution tubes in 96-tube array.*
- (i) *Reagent reservoirs.*
- (j) *Ice bucket and ice.*
- (k) *Vortex mixer.*

- (l) *Sealing tape.*—Millipore; Cat. No. MATA HCL00.
- (m) *Volumetric flask.*—1 L.
- (n) *–80°C freezer.*
- (o) *Refrigerator.*

For traditional scintillation counter only:

- (p) *MultiScreen punch device.*—Millipore; Cat. No. MAMP 096 08.
 - (q) *MultiScreen disposable punch tips.*—Millipore; Cat. No. MADP 196 10.
 - (r) *MultiScreen punch kit B for 4 mL vials.*—Millipore; Cat. No. MAPK 896 0B.
 - (s) *Scintillation vials.*—4 mL.
- For sample extraction:
- (t) *Pipets.*
 - (u) *Centrifuge tubes.*—15 mL, conical, plastic.
 - (v) *Vacuum pump or house vacuum.*
 - (w) *pH meter or pH paper.*
 - (x) *Hot plate.*
 - (y) *Graduated centrifuge tubes.*—15 mL.
 - (z) *Centrifuge and rotor for 15 mL tubes.*

C. Reagents

- (a) [^3H] STX.—0.1 mCi/mL, ≥ 10 Ci/mmol, $\geq 90\%$ radiochemical purity (American Radiolabeled Chemicals, St. Louis, MO, USA, or International Isotopes Clearinghouse, Leawood, KS, USA).
 - (b) STX diHCl.—NIST RM 8642 (www.nist.gov).
 - (c) *3-Morpholinopropanesulfonic acid (MOPS).*—Sigma (St. Louis, MO, USA; Cat. No. M3183-500G), or equivalent.
 - (d) *Choline chloride.*—Sigma (Cat. No. C7527-500G), or equivalent.
 - (e) *Rat brain membrane preparation.*—Appendix 1 [*J. AOAC Int.* (future issue)].
- For traditional counter:
- (f) *Scintiverse BD liquid scintillation cocktail.*—Fisher Scientific (Waltham, MA, USA; Cat. No. SX-18), or equivalent.
- For microplate counter:
- (g) *Optiphase liquid scintillation cocktail.*—PerkinElmer Life Sciences (Downers Grove, IL, USA; Cat. No. 1200-139), or equivalent.
- For sample extraction:
- (h) *Hydrochloric acid (HCl).*—1.0 and 0.1 M.
 - (i) *Sodium hydroxide.*—0.1 M.
 - (j) *Water.*—Distilled or deionized (18 $\mu\Omega$).

D. Sample Extraction

Accurately weigh 5.0 g tissue homogenate into a tared 15 mL conical tube. Add 5.0 mL of 0.1 M HCl, vortex, and check pH. If necessary, adjust pH to 3.0–4.0 as determined by a pH meter or pH paper. To lower pH, add 1 M HCl dropwise with mixing; to raise pH, add 0.1 M NaOH dropwise with mixing to prevent local alkalization and consequent destruction of toxin. Place the tube in a beaker of boiling water on hot plate for 5 min with the caps loosened. Remove and cool to room temperature. Check pH and adjust cooled mixture to pH 3.0–4.0 as described above. Transfer entire contents to graduated centrifuge tube and dilute volumetrically to 10 mL. Gently stir contents to homogeneity and allow to settle until portion of supernatant is translucent and can be decanted free of solid particles. Pour approximately 5 to 7 mL of the translucent supernatant into a centrifuge tube. Centrifuge at $3000 \times g$ for 10 min. Retain clarified supernatant and transfer to a clean centrifuge tube. Store extracts at -20°C until tested in

Table 2011.27A. Receptor binding assay results on individual samples (values are in µg STX diHCl equiv./kg shellfish tissue); summary statistics on all samples; summary statistics excluding Laboratory 9

Assay	No.	ID	Sample									All labs				Labs 1-8			
			1	2	3	4	5	6	7	8	9	Mean	S _R	RSD _R , %	HorRat	Mean	S _R	RSD _R , %	HorRat
Day 1	1	MLV05	370	610	620	410	690	1070	630	660	330	599	222	37.1	2.2	633	212	33.5	2.0
	2	MLV06	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	422	28.6	1.9	1549	386	24.9	1.7
Day 2	3	MLV08	80	190	140	90	130	160	230	220	100	149	55	37.2	1.8	155	56	36.0	1.7
	4	MLV09	860	680	950	870	980	1120	1460	820	590	926	255	27.5	1.7	968	237	24.5	1.5
Day 3	5	MLV12	180 ^a	200	200	150	150	100	150	290	100	168	62	37.2	1.8	177	60	34.1	1.7
	6	MLV14	950	940	1060	1130	1040	750	1460	1320	810	1051	228	21.7	1.4	1081	224	20.7	1.3
Day 3	7	MLV16	660	930	1080	870	840	1320	1490	2420 ^b	490	960	329	34.3	2.1	1027	291	28.3	1.8
	8	MLV01	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	818	48.6	3.3	1829	739	40.4	2.8
Day 3	9	MVL02	830	1180	1130	1150	1130	1780	1340	980	690	1134	311	27.4	1.8	1190	281	23.6	1.5
	10	MLV04	2440	2840	2910	1740	2150	1810	2690	2490	1210	2253	572	25.4	1.8	2384	446	18.7	1.3
Day 3	11	MLV07	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	345	23.9	1.6	1520	279	18.3	1.2
	12	MLV09	810	1190	1130	810	1630	1390	1880	1120	870	1203	372	30.9	2.0	1245	375	30.1	2.0
Day 3	13	MLV11	270	370	480	340	640	490	240	600	110	393	174	44.3	2.4	429	148	34.4	1.9
	14	MLV13	400	1240 ^b	560	450	650	530	500	440	200	466	133	28.5	1.6	504	85	16.8	1.0
Day 3	15	MLV03	330	270	410	180	590	680	370	1570 ^b	90	365	197	54.0	2.9	404	176	43.5	2.4
	16	MLV05	580	670	250	430	910	700	860	940	300	627	257	41.1	2.4	668	242	36.2	2.1
Day 3	17	MLV06	1290	1520	1460	970	1800	2520	1470	870	1250	1461	488	33.4	2.2	1488	515	34.6	2.3
	18	MLV07	1010	1600	1390	1000	1720	1860	1520	2150	890	1460	429	29.4	2.0	1531	397	26.0	1.7
Day 3	19	MLV10	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	570	24.0	1.7	2443	569	23.3	1.7
	20	MLV11	430	350	460	280	550	620	1149 ^b	410	250	419	127	30.2	1.7	443	115	26.0	1.4
Day 3	21	MLV15	ND ^c	ND	ND	ND	ND	ND	ND	180	ND	—	—	—	—	—	—	—	—
	Avg. RSD _R																		
	Avg. HorRat												33.2	2.0					28.7

^a CV 41%, not used in calculations.
^b Outlier; not used in calculations.
^c ND = Not detected.

Table 2011.27B. Summary statistics on blind duplicates, run in separate assays (values are in µg STX diHCl equiv./kg)

Lab	MLV05		MLV06		MLV07		MLV09		MLV11		Avg.
	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	
1	370	580	1100	1290	1260	1010	860	810	270	430	
2	610	670	1340	1520	1540	1530	680	1190	370	350	
3	620	250	1320	1460	1220	1390	950	1130	480	401	
4	410	430	1440	970	1980	1000	870	810	340	280	
5	690	910	1260	1790	1760	1720	980	1630	640	550	
6	1070	700	1720	2520	1530	1860	1120	1390	490	620	
7	630	880	2090	1240	1750	1150	1460	1830	230 ^a	1149 ^a	
8	660	940	2130	870	1210	2150	820	1120	600	410	
9	330	300	890	1250	840	890	590	870	110	250	
Avg.		614		1453		1433		1062		416	
S _r		169		432		366		247		83	
S _R		239		444		387		338		152	
RSD _r %		27.5		29.4		25.5		23.3		20.0	25.1
RSD _R %		38.9		30.2		27.0		31.9		36.5	32.9
HorRat		2.3		2.0		1.8		2.0		2.0	2.0

^a Outlier; not used in calculation.

receptor assay.

E. Preparation of Stock Solutions and Standards

(a) *Assay buffer*.—100 mM MOPS/100 mM choline chloride, pH 7.4. Weigh out 20.9 g MOPS and 13.96 g choline chloride and add to 900 mL dH₂O. Adjust pH to 7.4 with NaOH while stirring and bring to a final volume of 1 L with dH₂O. Store at 4°C.

(b) *Radioligand solution*.—Calculate the concentration of [³H] STX stock provided by the supplier, which may vary between lots. Suppliers generally provide the specific activity in Ci/mmol (generally 10–30 Ci/mmol) and activity in mCi/mL (0.05–0.1 mCi/mL), from which the molar concentration can be calculated. Prepare 4 mL of a 15 nM working stock of [³H] STX fresh daily in 100 mM MOPS/100 mM choline chloride buffer. This will provide sufficient volume for one 96-well plate at an in-well concentration of 2.5 nM. Measure total counts of each working stock prior to running an assay: add 35 µL of the working stock [³H] STX in buffer to a liquid scintillation vial with 4 mL scintillant and count on a traditional liquid scintillation counter. This is done to confirm correct dilution prior to running the assay. Depending on the efficiency of the scintillation counter used, the corresponding CPM will vary, but should be consistent day-to-day and within 15% of the expected value.

(c) *Unlabeled STX standard working solution*.—The STX diHCl standard is provided at a concentration of 268.8 µM (100 µg/mL). A “bulk” standard curve can be made up in advance and stored at 4°C for up to 1 month. The use of a bulk standard curve minimizes the pipetting needed for setting up an assay routinely and improves day-to-day repeatability. Make up 3 mM HCl (e.g., from a 3 M stock, 50 µL in 50 mL), then perform the serial dilutions (see Table 2011.27F) of NIST RM 8642 STX diHCl (100 µg/mL = 268.8 µM) to make up the standard curve in 3 mM HCl. These standard stock solutions will be diluted 1/6 in the assay to yield the designated in-assay concentrations (see Table 2011.27F).

(d) *Interassay calibration standard (QC check)*.—Prepare a

reference standard containing 1.8×10^{-8} M STX standard (3.0×10^{-9} M STX in assay) in advance in 3 mM HCl and keep frozen (–80°C) in 1 mL aliquots for long-term storage. Aliquots should be thawed and stored at 4°C for routine use (stable up to 1 month) and analyzed in each assay. This serves as a QC check and confirms day-to-day performance of the assay.

(e) *Rat brain membrane preparation*.—Prepare rat brain membrane preparation in bulk [Appendix I; J. AOAC Int. (future issue)] and store at –80°C until used in the assay. Thaw an aliquot of rat brain membrane preparation on ice. Dilute membrane preparation with cold (4°C) 100 mM MOPS/100 mM choline chloride, pH 7.4, to yield a working stock with a protein concentration of 1.0 mg/mL (this will be diluted in the assay plate to 0.5 mg/mL in-well concentration). Vortex vigorously to achieve a visibly homogeneous suspension. Keep the diluted membrane preparation on ice until ready to use.

F. Performing the Assay

(a) *Plate setup*.—When possible, use a multichannel pipet to minimize pipetting effort and increase consistency. Standard curve, QC check, and sample extracts are run in triplicate wells. Multiple dilutions of sample extracts should be analyzed in order to obtain a value that falls between 0.2–0.7 B/B₀ on the standard curve for quantification. For ease of analysis, it is convenient to use a standard plate layout that maximizes the number of samples and standards that can be analyzed on one plate. For shellfish extracts, a minimum dilution of 1:10 is used, which minimizes potential matrix effects, while still providing an LOQ of approximately 126 µg/kg shellfish (see Table 2011.27G).

(b) *Addition of samples and standards*.—Add in the following order to each of the 96 wells: 35 µL assay buffer; 35 µL STX standard, QC check, or sample extract; 35 µL [³H] STX; 105 µL membrane preparation. The assay buffer is added first in order to wet the filter membrane. It is critical to continuously mix the membrane preparation by careful up-and-down pipetting immediately prior to

Table 2011.27C. Performance of individual laboratories on blind duplicates (values are in µg STX diHCl equiv./kg)

Laboratory	ID	Day 1	Day 2	Mean	s _p	RSD _r , %
1	MLV05	370	580	475	148	31.3
	MLV06	1100	1290	1195	134	11.2
	MLV07	1260	1010	1135	177	15.6
	MLV09	860	810	835	35	4.2
	MLV11	270	430	350	113	32.3
Avg.						18.9
2	MLV05	605	670	638	46	7.2
	MLV06	1340	1520	1430	127	8.9
	MLV07	1540	1530	1535	7	0.5
	MLV09	680	1190	935	361	38.6
	MLV11	370	350	360	14	3.9
Avg.						11.8
3	MLV05	620	250	435	262	60.1
	MLV06	1320	1460	1390	99	7.1
	MLV07	1220	1303	1262	59	4.7
	MLV09	950	1130	1040	127	12.2
	MLV11	480	460	470	14	3.0
Avg.						17.4
4	MLV05	410	430	420	14	3.4
	MLV06	1440	970	1205	332	27.6
	MLV07	1980	1000	1490	693	46.5
	MLV09	870	810	840	42	5.1
	MLV11	340	280	310	42	13.7
Avg.						19.2
5	MLV05	690	910	800	156	19.4
	MLV06	1260	1790	1525	375	24.6
	MLV07	1760	1720	1740	28	1.6
	MLV09	980	1630	1305	460	35.2
	MLV11	640	550	595	64	10.7
Avg.						18.3
6	MLV05	1070	700	885	262	29.6
	MLV06	1720	2520	2120	566	26.7
	MLV07	1530	1860	1695	233	13.8
	MLV09	1120	1390	1255	191	15.2
	MLV11	490	620	555	92	16.6
Avg.						20.4
7	MLV05	630	880	755	177	23.4
	MLV06	2090	1240	1665	601	36.1
	MLV07	1750	1150	1450	424	29.3
	MLV09	1460	1830	1645	262	15.9
	MLV11	230 ^a	1150 ^a			
Avg.						26.2
8	MLV05	660	940	800	198	24.7
	MLV06	2130	870	1500	891	59.4
	MLV07	1210	2150	1680	665	39.6
	MLV09	820	1120	970	212	21.9
	MLV11	600	410	505	134	26.6
Avg.						34.4
9	MLV05	330	300	315	21	6.7
	MLV06	890	1250	1070	255	23.8
	MLV07	840	890	865	35	4.1
	MLV09	590	870	730	198	27.1
	MLV11	110	250	180	99	55.0
Avg.						23.3
Overall avg.						22.2

^a Outlier; not used in calculations.

dispensing into the 96-well plate to maintain an even suspension across the entire plate. Cover and incubate plate at 4°C for 1 h.

(c) *Assay filtration.*—Attach the vacuum manifold to the vacuum pump with an in-line side arm flask to catch filtrate from the plate filtration process. Set the vacuum pressure gauge on the pump or vacuum manifold to 4–8" Hg (135–270 millibar), as specified in the instructions provided with the filtration plates. Place the 96-well plate on the vacuum manifold. Fill empty wells with 200 µL MOPS/choline chloride buffer to ensure even vacuum pressure and filtration across the plate. Turn on vacuum. Optimum vacuum will pull the wells to dryness in 2–5 s. Pull contents of all wells through until all liquid is removed. (*Note:* Too low a vacuum will result in slow well clearance, but too high will result in an airlock and no well clearance.) With vacuum pump running, quickly rinse each well twice with 200 µL ice cold MOPS/choline chloride buffer using multichannel pipet. Maintain vacuum until liquid is removed.

(d) *Preparation of the assay for counting.*—Remove the plastic bottom from the plate. Blot the bottom once on absorbent toweling.

(1) *For counting in microplate scintillation counter.*—Place the microplate in a counting cassette. Seal the bottom of the 96-well plate with sealing tape. Add 50 µL Optiphase scintillation cocktail per well using multichannel pipet. Seal the top of the plate with sealing tape. Allow to incubate 30 min at room temperature. Place the plate in a counting cassette and count in a microplate scintillation counter for 1 min/well.

(2) *For counting in traditional scintillation counter.*—Place the microplate in the MultiScreen punch system apparatus. Place the disposable punch tips on top of the microplate. Punch the filters from the wells into scintillation vials and fill with 4 mL scintillation cocktail (Scintiverse or equivalent). Place caps on the vials and vortex. Allow vials to sit overnight in the dark, then count using a tritium window in a traditional scintillation counter.

G. Analysis of Data

For assays performed using the traditional counter, curve fitting is performed using a four-parameter logistic fit, also known as a sigmoidal dose response curve (variable slope; *see* Figure 2011.27), or Hill equation:

$$y = \min + \frac{\max - \min}{1 + 10^{(x - \log IC_{50}) \text{Hill slope}}}$$

where max is the top plateau representing maximum binding in CPM in the absence of competing nonradiolabeled STX, also known as B₀; min is the bottom plateau, equal to nonspecific binding (in CPM) in the presence of saturating nonradiolabeled toxin; IC₅₀ is the inhibitory concentration at which CPM are 50% of max-min (dashed lines; Figure 2011.27); Hill slope is the slope of the curve; x axis is the log concentration of STX; and y axis is total ligand binding in CPM (here represented as B/B₀, or bound/max bound). A curve fitting package such as Prism (Graph Pad Software, Inc.) is recommended. For the microplate counter users, receptor assay applications provided by the manufacturer may be used (e.g., MultiCalc; PerkinElmer Wallac, Gaithersburg, MD, USA).

(a) *Sample quantification.*—Sample quantification is carried out only on dilutions that fall within B/B₀ of 0.2–0.7, where B represents the bound [³H]STX (in CPM) in the sample and B₀ represents the max bound [³H]STX (in CPM). Where more than one dilution falls within B/B₀ of 0.2–0.7 on the curve, all sample wells corresponding to these dilutions are used to calculate sample concentration. Sample concentration is calculated in µg STX diHCl

Table 2011.27D. Calibration curve and QC check parameters in three receptor binding assays performed in nine participant laboratories

Lab	Assay day	Slope	IC ₅₀ ^a , nM	QC, nM	Reference, CPM	IC ₇₀ ^a , nM	Standards where RSD >30%; action	Curve fitting software	Scintillation counter	Manual/microplate
1	1	-0.9	1.9	2.4	720	0.90	None	Prism v 3.02	Packard Top Count	Microplate
	2	-1.0	2.0	2.6	733	0.96	None			
	3	-1.1	2.1	3.2	1038	0.92	None			
2	1	-1.1	1.8	3.8	1160	0.66	3 nM; 1 well removed	Prism v 5.0	Packard Top Count	Microplate
	2	-1.2	2.2	3.9	1260	0.85	None			
	3	-1.0	1.6	3.2	1262	0.46	3 nM, 1 nM removed			
3	1	-1.0	2.0	2.3	2529	0.41	First column removed	Prism v 5.0	Wallac Microbeta	Microplate
	2	-0.9	2.0	2.5	1463	0.92	1000 nM; 1 well removed			
	3	1.0	1.6	2.8	2088	0.80	None			
4	1	-0.9	1.7	3.4	1125	0.61	None	Prism v 3.03	PerkinElmer Tricarb	Manual
	2	-1.2	1.7	3.2 ^a	1611	0.77	None			
	3	-0.9	1.2	2.9	1324	0.45	30 nM 35%; 1 well removed			
5	1	-0.9	1.4	3.3	1566	0.64	1.0 nM; 1 well removed	MultiCalc	Wallac Microbeta	Microplate
	2	-1.2	1.8	3.6	1528	1.05	0.1 nM and 30 nM; 1 well removed			
	3	-1.2	1.8	2.9	1052	0.67	None			
6	1	-1.1	2.6	3.0	670	1.15	None	Prism v 4.0	Wallac Microbeta	Microplate
	2	-1.0	2.0	4.0 ^b	1124	1.08	None			
	3	-1.1	3.4	6.5 ^b	1030	2.04 ^c	None			
7	1	-0.8	1.0	2.8 ^a	919	0.33	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	1.6	2.7	619	0.70	None			
	3	-0.9	2.1	3.2 ^a	693	0.82	None			
8	1	-1.2	1.7	3.7	1146	0.86	None	Prism	Wallac Microbeta	Microplate
	2	-1.1	1.4	1.5 ^b	1095	0.78	None			
	3	-1.1	2.4	2.3	886	1.04	None			
9	1	-1.0	2.2	4.0 ^b	1363	0.97	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	2.0	3.2	1380	0.85	100 nM 33%; left in			
	3	-1.0	2.1	3.7	1532	0.92	None			

^a One well removed.

^b Outside of specifications.

^c Outlier by Grubbs test.

Table 2011.27E. Results of the receptor binding assay (RBA), mouse bioassay (MBA), and HPLC analyses of 21 shellfish extracts, sorted by mouse bioassay value (all values are in µg STX diHCl equiv./kg shellfish tissue; results in bold indicate toxicity above the 800 µg STX diHCl equiv./kg regulatory limit; all other results indicate toxicity below the regulatory limit)

Sample	Lab 1	Lab 2	Lab 3	Lab 4	Lab 5	Lab 6	Lab 7	Lab 8	Lab 9	RBA, avg.	HPLC	MBA
21	ND ^a	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
5	180	200	200	150	150	100	150	290	100	168	108	ND
15	330	270	410	180	590	680	370	1570^b	90	365	196	182
13	270	370	480	340	640	290	240	600	110	371	236	299
20	430	350	460	280	550	490	1150^b	410	250	403	236	299
14	400	1240^b	560	450	650	530	500	440	200	466	625	343
1	370	610	620	410	690	1070^b	630^b	660	330	599	413	387
16	580	670	250	430	910	700	860^b	940^b	300	627	413	387
3	80	190	140	90	130	160	230	220	100	149	341	405
6	950	940	1060	1130	1040	750	1460	1320	810	1051	618	485
7	660	930	1080	870	840	1320	1490	2420	490	960	685	528
2	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	931	595
17	1290	1520	1460	970	1800	2520	1470	870	1250	1460	931	595
4	860	680	950	870	980	1120	1460	820	590	926	1070	653
12	810	1190	1130	810	1630	1390	1880	1120	870	1203	1070	653
11	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	965	714
18	1010	1600	1390	1000	1720	1860	1520	2150	890	1452	965	714
8	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	894	752
9	830	1180	1130	1150	1130	1780	1340	980	690	1134	802	792
19	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	2000	1027
10	2440	2840	2910	1740	2150	1800	2690	2490	1210	2252	1890	1080

^a ND = Not detected.

^b Outlier; not used in average calculation.

equiv./kg shellfish, using the following formulas:

$$(\text{nM STX equiv.}) \times (\text{sample dilution}) \times \frac{(210 \mu\text{L total volume})}{35 \mu\text{L sample}} = \text{nM STX equiv. in extract}$$

$$(\text{nM STX diHCl equiv. in extract}) \times \frac{1 \text{ L}}{1000 \text{ mL}} \times \frac{372 \text{ ng}}{\text{nmol}} \times \frac{1 \mu\text{g}}{1000 \text{ ng}} = \mu\text{g STX diHCl equiv./mL}$$

$$\mu\text{g STX diHCl equiv./mL} \times \frac{\text{mL extract}}{\text{g shellfish}} \times \frac{1000 \text{ g}}{\text{kg}} = \mu\text{g STX diHCl equiv./kg}$$

H. Assay Performance Standards

The following criteria must be met for assay acceptance:

(a) For a ligand that specifically binds at one receptor site, the slope of the resulting competition curve should theoretically be -1.0. If the slope of the curve for a given assay is outside of the acceptable range of -0.8 to -1.2, linearity of the assay will be compromised and quantification of the unknowns will be incorrect.

(b) RSDs of triplicate CPMs for standards should be below 30% as variability may affect the slope calculation and thereby quantification of samples.

(c) If the IC₅₀ is out of the acceptable range (2.0 nM ± 30%)

then the assay should be considered suspect and rerun, as a shift in the curve will result in over- or underestimation of sample concentrations.

(d) QC check should be 3 nM STX ± 30% (in-well concentration).

Table 2011.27F. Dilution series to prepare bulk solutions for standard curve

	Stock, M	In-assay, M
100 µL 268.8 µM STX + 4.38 mL 0.003 M HCl	6 × 10 ⁻⁶	1 × 10 ⁻⁶
500 µL 6 × 10 ⁻⁶ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁷	1 × 10 ⁻⁷
1.5 mL 6 × 10 ⁻⁷ M + 3.5 mL 0.003 M HCl	1.8 × 10 ⁻⁷	3 × 10 ⁻⁸
500 µL 6 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁸	1 × 10 ⁻⁸
500 µL 1.8 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	1.8 × 10 ⁻⁸	3 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁸ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁹	1 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁹ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻¹⁰	1 × 10 ⁻¹⁰
5 mL 0.003 M HCl	0	Reference

Table 2011.27G. Recommended microplate layout for ease of handling triplicate wells of standard curve, QC check sample, and unknown samples; each sample is run at three dilutions (1:10, 1:50, 1:200); standard curve is run in columns 1–3 (values are in M STX)^a

Microplate row	Microplate column											
	1	2	3	4	5	6	7	8	9	10	11	12
A	10 ⁻⁶	10 ⁻⁶	10 ⁻⁶	QC	QC	QC	U3 1:50	U3 1:50	U3 1:50	U6 1:10	U6 1:10	U6 1:10
B	10 ⁻⁷	10 ⁻⁷	10 ⁻⁷	U1 1:10	U1 1:10	U1 1:10	U3 1:200	U3 1:200	U3 1:200	U6 1:50	U6 1:50	U6 1:50
C	3 × 10 ⁻⁸	3 × 10 ⁻⁸	3 × 10 ⁻⁸	U1 1:50	U1 1:50	U1 1:50	U4 1:10	U4 1:10	U4 1:10	U6 1:200	U6 1:200	U6 1:200
D	10 ⁻⁸	10 ⁻⁸	10 ⁻⁸	U1 1:200	U1 1:200	U1 1:200	U4 1:50	U4 1:50	U4 1:50	U7 1:10	U7 1:10	U7 1:10
E	3 × 10 ⁻⁹	3 × 10 ⁻⁹	3 × 10 ⁻⁹	U2 1:10	U2 1:10	U2 1:10	U4 1:200	U4 1:200	U 1:200	U7 1:50	U7 1:50	U7 1:50
F	10 ⁻⁹	10 ⁻⁹	10 ⁻⁹	U2 1:50	U2 1:50	U2 1:50	U5 1:10	U5 1:10	U5 1:10	U7 1:200	U7 1:200	U7 1:200
G	10 ⁻¹⁰	10 ⁻¹⁰	10 ⁻¹⁰	U2 1:200	U2 1:200	U2 1:200	U5 1:50	U5 1:50	U5 1:50			
H	REF	REF	REF	U3 1:10	U3 1:10	U3 1:10	U5 1:200	U5 1:200	U5 1:200			

^a REF = Reference; QC = quality control check; U = unknown sample. [Note: The same standard curve may be used for multiple plates (i.e., 11 samples can be run on subsequent plates in a series if the standard curve is not included).]

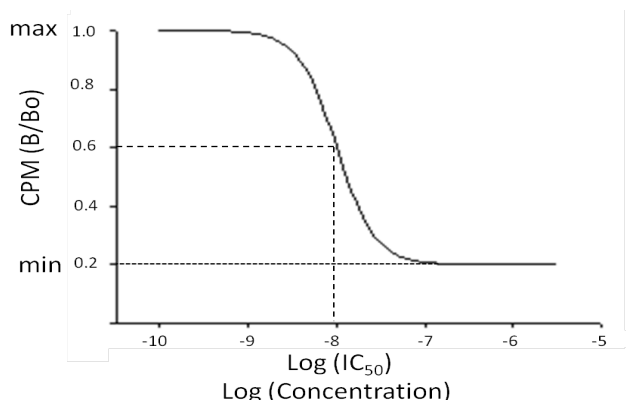


Figure 2011.27. Sigmoidal dose response curve. Dashed lines indicate log IC50.

Assays with a QC check sample out of specifications should trigger a check of the IC₅₀ value.

The following criteria must be met for acceptability of a sample measurement:

(a) Sample quantification should be done only on dilutions that fall within B/B₀ of 0.2–0.7. In the event that all sample dilutions fall below B/B₀ 0.2 (i.e., concentration is too high), further dilutions must be made and the sample reanalyzed. In the event that the sample concentration is too low to be quantified (i.e., B/B₀ > 0.7), the sample is reported as below LOD. If more than one dilution falls on the linear part of the curve, an average value calculated from all dilutions should be used. If there is disagreement between different dilutions in final concentration reported, check for error in the sample dilution process.

(b) RSD of the sample CPMs should be ≤30%.

Reference: *J. AOAC Int.* (future issue)

Single-Laboratory Validation of the Microplate Receptor Binding Assay for Paralytic Shellfish Toxins in Shellfish

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A single-laboratory validation (SLV) study was conducted for the microplate receptor binding assay (RBA) for paralytic shellfish poisoning (PSP) toxins in shellfish. The basis of the assay is the competition between [³H]saxitoxin (STX) and STX in a standard or sample for binding to the voltage dependent sodium channel. A calibration curve is generated by the addition of 0.01–1000 nM STX, which results in the concentration dependent decrease in [³H]STX-receptor complexes formed and serves to quantify STX in unknown samples. This study established the LOQ, linearity, recovery, accuracy, and precision of the assay for determining PSP toxicity in shellfish extracts, as performed by a single analyst on multiple days. The standard curve obtained on 5 independent days resulted in a half-maximal inhibition (IC₅₀) of 2.3 nM STX ± 0.3 (RSD = 10.8%) with a slope of 0.96 ± 0.06 (RSD = 6.3%) and a dynamic range of 1.2–10.0 nM. The LOQ was 5.3 µg STX equivalents/100 g shellfish. Linearity, established by quantification of three levels of purified STX (1.5, 3, and 6 nM), yielded an r² of 0.97. Recovery from mussels spiked with three levels (40, 80, and 120 µg STX/100 g) averaged 121%. Repeatability (RSD_r), determined on six naturally contaminated shellfish samples on 5 independent days, was 17.7%. A method comparison with the AOAC mouse bioassay yielded r² = 0.98 (slope = 1.29) in the SLV study. The effects of the extraction method on RBA-based toxicity values were assessed on shellfish extracted for PSP toxins using the AOAC mouse bioassay method (0.1 M HCl) compared to that for the precolumn oxidation HPLC method (0.1% acetic acid). The two extraction methods showed linear correlation (r² = 0.99), with the HCl extraction method yielding slightly higher toxicity values (slope = 1.23). A similar relationship was

observed between HPLC quantification of the HCl- and acetic acid-extracted samples (r² = 0.98, slope 1.19). The RBA also had excellent linear correlation with HPLC analyses (r² = 0.98 for HCl, r² = 0.99 for acetic acid), but gave somewhat higher values than HPLC using either extraction method (slope = 1.39 for HCl extracts, slope = 1.32 for acetic acid). Overall, the excellent linear correlations with the both mouse bioassay and HPLC method and sufficient interassay repeatability suggest that the RBA can be effective as a high throughput screen for estimating PSP toxicity in shellfish.

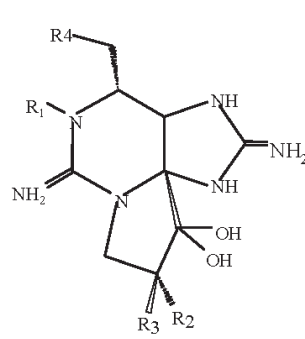
Paralytic shellfish poisoning (PSP) is a seafood intoxication caused by the consumption of shellfish tainted with saxitoxins (STXs) produced by certain species of harmful algae. Saxitoxins are a suite of heterocyclic guanidinium toxins, of which currently more than 21 congeners are known (Figure 1). These congeners occur in varying proportions in the dinoflagellates that produce them and are further metabolized in shellfish that accumulate them, making analytical determination of PSP toxins in shellfish complex. The long-standing regulatory method for PSP toxins is the AOAC mouse bioassay (1), with a regulatory limit of 80 µg/100 g shellfish generally applied. Increasing resistance to whole animal testing has driven the need to develop alternative methods suitable for use in a high throughput monitoring or regulatory setting. In the past decade, several alternatives to the mouse bioassay have been developed and validated to various degrees. The precolumn oxidation HPLC method (2) has received First Action approval by AOAC as an Official Method for PSP (2005.06; 3) and has been accepted into the European Food Hygiene Regulations as an alternative to the mouse bioassay and further refined to optimize its use in the United Kingdom Official Control monitoring of PSP toxins in mussels (4). However, although the HPLC method performs well quantitatively, it is quite time consuming for high throughput screening needed by many monitoring programs. A qualitative lateral flow antibody test for PSP toxins with a detection limit of 40 µg/100 g, developed by

Jellett Rapid Testing Ltd (Chester Basin, NS, Canada), has been approved in the United States by the Interstate Shellfish Sanitation Conference and the U.S. Food and Drug Administration (FDA) as a screening method. This method performed well in a comparison study with the mouse bioassay, with a false-positive rate of 6% and a false-negative rate of <0.1% (5), but it has not been put through a full AOAC collaborative trial, and does not provide quantitative analysis. To date, a suitable quantitative, high throughput alternative to the mouse bioassay has not been validated through the AOAC Official Methods Program. The current study establishes the single laboratory performance characteristics of the microplate receptor binding assay (RBA) for PSP toxins in shellfish and identifies it as a candidate for fulfilling the requirements of high throughput, quantitative analysis that measures a composite toxic potency in a manner analogous to the mouse bioassay.

STX elicit their paralytic effects by binding to site 1 on the voltage dependent sodium channel, thereby blocking the transmission of neuronal and muscular action potentials. Because all STX congeners bind to site 1 with affinities proportional to their mouse intraperitoneal (IP) toxicity (6), a receptor binding competition assay can be used to measure the integrated toxic potency of STX congeners in a sample, independent of which toxin congeners are present. Moreover, any toxin metabolites originating in the shellfish matrix will also be detected by the assay according to their affinity for the sodium channel receptor. In this binding competition assay, [³H]STX competes with unlabeled STX and/or its derivatives for a finite number of available receptor sites in a rat brain membrane preparation. Following establishment of binding equilibrium, unbound [³H]STX is removed by filtration and

bound [³H]STX is quantified by liquid scintillation counting. The percent reduction in [³H]STX binding in the presence of unlabeled toxin is directly proportional to the amount of unlabeled toxin present. A standard curve is established using increasing concentrations of unlabeled STX, and the concentration of PSP toxins in an unknown sample is quantified using this standard curve.

The assay tested in this single laboratory trial is a modification of the method of Doucette et al. (7) to a 96-well microplate format described by Van Dolah et al. (8). Application of microplate scintillation counting to the PSP assay was first reported by Powell and Doucette (9), who applied it to phytoplankton analysis. The use of the microplate format, in conjunction with microplate scintillation counting, makes the assay suitable for use in a high throughput monitoring or regulatory setting. Several versions of the PSP receptor binding assay have undergone method comparisons in different laboratories with favorable correlations to the mouse bioassay and/or other assays for PSP toxins in shellfish. Suarez-Isla and Valez (10) showed excellent linear correlation ($r^2 = 0.97$) between the RBA and mouse bioassay of 41 shellfish extracts between 40 and 10 000 μg STX equivalents/100 g. Llewellyn et al. (11) found that the sodium channel receptor assay compared well to three other methods of analysis for PSP toxins in shellfish (HPLC, mouse bioassay, and N2A cytotoxicity assay). Ruberu et al. (12) optimized the microplate format assay for use in the Packard Top Count microplate scintillation counter (a single channel counter; GMI, Inc., Ramsey, MN), compared results with the same assay performed on the Wallac microplate counter (a two-channel coincidence counter; Perkin Elmer Wallace, Gaithersburg, MD), and provided further correlation data with



		R1	R2	R3	R4	MU/ μmol
Carbamate	STX	H	H	H	OCONH ₂	2483
	Neo STX	OH	H	H	OCONH ₂	2295
	GTX1	OH	OSO ₃ -	H	OCONH ₂	2468
	GTX2	H	OSO ₃ -	H	OCONH ₂	892
	GTX3	H	H	OSO ₃ -	OCONH ₂	1584
	GTX4	OH	H	OSO ₃ -	OCONH ₂	1803
Sulfocarbamoyl	GTX5 (B1)	H	H	H	OCONHSO ₃ -	160
	GTX6 (B2)	OH	H	H	OCONHSO ₃ -	-
	C1	H	OSO ₃ -	H	OCONHSO ₃ -	15
	C2	H	H	OSO ₃ -	OCONHSO ₃ -	239
	C3	OH	OSO ₃ -	H	OCONHSO ₃ -	33
C4	OH	H	OSO ₃ -	OCONHSO ₃ -	143	
Decarbamoyl	dcSTX	H	H	H	OH	1274
	dcNeoSTX	OH	H	H	OH	-
	dcGTX1	OH	OSO ₃ -	H	OH	-
	dcGTX2	H	OSO ₃ -	H	OH	1617
	dcGTX3	H	H	OSO ₃ -	OH	1872
dcGTX4	OH	H	OSO ₃ -	OH	-	
Deoxydecarbamoyl	doSTX	H	H	H	H	-
	doGTX2	H	H	OSO ₃ -	H	-
	doGTX3	H	OSO ₃ -	H	H	-

Figure 1. Structures and toxic potency of 21 saxitoxin congeners. Toxic potency is listed as mouse units (MU)/ μmole , where a mouse unit is defined as the minimum amount required to kill a 20 g mouse in 15 min when administered by IP injection. The table is modified from ref. 15.

	1	2	3	4	5	6	7	8	9	10	11	12
A	10 ⁻⁶	10 ⁻⁶	10 ⁻⁶	REF	REF	REF	U3 1:10	U3 1:10	U3 1:10	U5 1:200	U5 1:200	U5 1:200
B	10 ⁻⁷	10 ⁻⁷	10 ⁻⁷	QC	QC	QC	U3 1:50	U3 1:50	U3 1:50	U6 1:10	U6 1:10	U6 1:10
C	3 x 10 ⁻⁸	3 x 10 ⁻⁸	3 x 10 ⁻⁸	U1 1:10	U1 1:10	U1 1:10	U3 1:200	U3 1:200	U3 1:200	U6 1:50	U6 1:50	U6 1:50
D	10 ⁻⁸	10 ⁻⁸	10 ⁻⁸	U1 1:50	U1 1:50	U1 1:50	U4 1:10	U4 1:10	U4 1:10	U6 1:200	U6 1:200	U6 1:200
E	3 x 10 ⁻⁹	3 x 10 ⁻⁹	3 x 10 ⁻⁹	U1 1:200	U1 1:200	U1 1:200	U4 1:50	U4 1:50	U4 1:50	U7 1:10	U7 1:10	U7 1:10
F	10 ⁻⁹	10 ⁻⁹	10 ⁻⁹	U2 1:10	U2 1:10	U2 1:10	U4 1:200	U4 1:200	U 1:200	U7 1:50	U7 1:50	U7 1:50
G	10 ⁻¹⁰	10 ⁻¹⁰	10 ⁻¹⁰	U2 1:50	U2 1:50	U2 1:50	U5 1:10	U5 1:10	U5 1:10	U7 1:200	U7 1:200	U7 1:200
H	10 ⁻¹¹	10 ⁻¹¹	10 ⁻¹¹	U2 1:200	U2 1:200	U2 1:200	U5 1:50	U5 1:50	U5 1:50			

U = unknown sample

Figure 2. Standardized plate layout recommended for the microplate RBA for PSP toxins in shellfish extracts. U = unknown sample.

the mouse bioassay. Usup et al. (13) utilized the microplate RBA method to compare predicted toxicity values in samples spiked with different STX congeners as assayed by the mouse bioassay and the RBA. Llewellyn (14) defined the competitive behavior of PSP toxin mixtures in receptor binding assays, using both the sodium channel and saxiphilin receptors, which explains their composite toxicity. However, none of these previous studies fully characterized assay performance according to AOAC single-laboratory validation (SLV) criteria that are the underpinning required for proceeding with an AOAC collaborative trial. Therefore, the current study was carried out to fulfill those requirements.

Experimental

Apparatus

- (a) *Microplate scintillation counter*.—Wallac Microbeta, GMI Inc. (Ramsey, MN).
- (b) *Microplate filtration manifold*.—Millipore (Bedford, MA).
- (c) *Hot plate*.—Fisher Scientific (Suwanee, GA).
- (d) *Countertop centrifuge*.—For 15 mL tubes, capable of 3000 × g (Fisher Scientific).
- (e) *Microtiter filter plates (96 well) with 1.0 μm pore size type FB glass fiber filter/0.65 μm pore size Duropore support membrane*.—Cat. No. MSFB N6B 50 (Millipore Corp., Billerica, MA).
- (f) *Microplate sealing tape*.—Cat. No. MATA HCL00 (Millipore Corp.).

(g) *Vortex mixer*.—Daigger Vortex Genie II (Daigger Scientific, Vernon Hills, IL).

(h) *Teflon/glass tissue homogenizer*.—Wheaton (Millville, NJ).

(i) *Polytron homogenizer*.—Brinkmann Instruments (Westbury, NY).

Reagents

- (a) *Hydrochloric acid (HCl)*.—0.1 M.
- (b) [³H]STX.—0.1 mCi/mL, ≥10 Ci/mmol, ≥90% radiochemical purity (International Isotopes Clearinghouse, Leawood, KS).
- (c) *STX diHCl*.—FDA reference standard (Office of Seafood, Laurel, MD) or National Research Council (NRC) of Canada Institute of Marine Biosciences (Halifax, NS, Canada).
- (d) *Assay buffer*.—75 mM HEPES [4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid; Cat. No. H9136]/140 mM NaCl, pH 7.5 (Sigma, St. Louis, MO).
- (e) *Liquid scintillation cocktail*.—Optiphase (PerkinElmer Life Sciences, Downers Grove, IL).

Preparation of Samples (0.1 M HCl Extraction)

Shellfish samples were shucked and homogenized according to the AOAC mouse bioassay protocol (1). For the HCl extraction method, 5.0 (±0.1) g of tissue homogenate was transferred to a tared 15 mL conical polypropylene centrifuge tube. A 5.0 mL volume of 0.1 M HCl was added, and the sample was mixed on a Vortex mixer. The pH was checked to

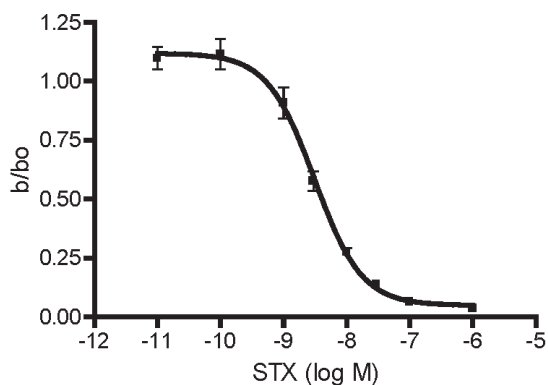


Figure 3. Average of five calibration curves obtained by one analyst in five independent assays on separate days. $IC_{50} = 2.23 \pm 0.23$ nM, slope = 0.96 ± 0.06 , error bars are \pm SD.

confirm it was between 3.0 and 4.0 in order to avoid alkalization and destruction of the toxin, and adjusted with 1 M HCl or 0.1 M NaOH as needed. Tubes were placed in a beaker of boiling water on a hot plate for 5 min with the caps loosened. Following removal from the boiling water bath, samples were allowed to cool to room temperature, and the pH was again confirmed to be between 3.0 and 4.0. The entire contents were then transferred to a graduated cylinder, diluted volumetrically to 10 mL, and centrifuged for 5 min at $1000 \times g$. The supernatant was transferred to a clean tube.

Preparation of Samples (Acetic Acid Extraction Method)

In a 50 mL plastic centrifuge tube, 5.0 ± 0.1 g homogenate was mixed with 3.0 mL 1% acetic acid on a vortex mixer. Tubes were capped loosely to avoid pressure buildup and placed in a boiling water bath for 5 min. Following removal from the water bath, samples were cooled in a beaker of cold water for 5 min, and then centrifuged for 10 min at $3000 \times g$. The supernatant was transferred to a 15 mL graduated conical test tube. A 3 mL amount of 1% acetic acid was added to the original tube with solid residue, mixed well on a vortex mixer, and centrifuged again for 10 min at $3000 \times g$. The second supernatant was combined with the first and diluted to 10 mL with water.

Preparation of Stock Solutions, Standards, and Reagents for Assay

(a) *Radioligand solution.*— $[^3H]$ STX stock is provided in 50 μ Ci ampules, 24 Ci/mmol, 0.1 mCi/mL (4.17 μ M). A 15 nM working stock of $[^3H]$ STX was prepared fresh daily in 75 mM HEPES/140 mM NaCl (for 2.5 nM final in-well concentration).

(b) *STX standard curve.*—FDA STX dihydrochloride reference standard (100 μ g/mL or 268.8 μ M) used to prepare a bulk standard curve made up in advance and stored at 4°C for up to 1 month. The stock standard curve was made consisted of eight concentrations of STX in 0.003 M HCl [6×10^{-6} , 6×10^{-7} , 1.8×10^{-7} , 6×10^{-8} , 1.8×10^{-8} , 6×10^{-9} , 6×10^{-10} , $6 \times$

Table 1. RBA measurements of calibration standards for assay linearity assessment (nM STX; $n = 5$)

Nominal	Mean	SD	RSD
1.5	1.7	0.16	10
3.0	3.0	0.52	17
6.0	6.0	0.34	6

10^{-11} , and 0.003 M only HCl (reference)], which when diluted 1:6 in the assay, resulted in a standard curve of 0.01 nM–1000 nM STX. The reference provided a measure of total $[^3H]$ STX binding in the absence of unlabeled STX.

(c) *Calibration standard (QC check).*—A reference standard containing 1.8×10^{-8} M STX standard (3.0×10^{-9} M STX in assay) was prepared in 0.003 M hydrochloric acid, aliquotted in 1 mL volumes, and stored at 4°C for routine use (stable up to 1 month). On the day of the assay, 200 μ L of each standard were pipetted into mini-dilution tubes for ease of pipetting into the microplate using an eight-channel pipettor.

(d) *Rat brain membrane homogenate.*—Cerebral cortices from 6-week-old male Holzman rats (Harlan Bioproducts, Indianapolis, IN) were homogenized on ice in a glass/Teflon tissue homogenizer in 75 mM HEPES/140 mM NaCl, pH 7.5, containing 0.1 mM PMSF (phenylmethanesulfonylfluoride; 12.5 mL/brain) at 385 rpm for 10 strokes. Pooled homogenates were centrifuged at $20\,000 \times g$ for 15 min at 4°C and the pellet was resuspended in HEPES buffer (12.5 mL/brain) and rehomogenized on ice using a Polytron homogenizer set at 70% power for 20 s to ensure a fine suspension. The brain homogenate was aliquotted 2 mL/tube in cryovials and stored at -80°C . The protein concentration of the brain homogenate was determined using the Micro bicinchoninic acid (BCA) Assay (Pierce, Rockford, IL). For each assay, an aliquot of brain homogenate was thawed on ice and diluted with ice cold 75 mM HEPES/150 mM NaCl, pH 7.5, to yield a final protein concentration of 0.5 mg/mL in the assay.

Table 2. Recovery of analyte from spiked samples (μ g STX equiv./100 g)

Nominal	Mean	SD	Measured RSD _r	Recovery, %
0	<dl ^a			
40	47	8.6	18.7	115
80	103.7	21.8	21	129
120	145.5	15.2	10.5	121

^a <dl = Less than LOQ (5 μ g STX equiv./100 g).

Table 3. Comparison of receptor binding assay (RBA; n = 5) with AOAC mouse bioassay (MBA) of naturally contaminated shellfish (µg STX equiv./100 g)

Sample	MBA	RBA mean	SD	RSD
LP1	340	438	74	17
LP2	534	715	96	13
LP3	1158	1533	329	21
LP4	65	91	7	9
LP5	350	608	150	25
LP6	462	518	114	22

Assay Procedure

(a) *Plate setup and incubation.*—A standardized plate layout was used for all assays (Figure 2). All standards, reference, QC check, and shellfish extracts were run in triplicate wells. For shellfish extracts, a standardized dilution series was run for each sample (1:10, 1:50, and 1:200), which ensured that at least one dilution would fall on the linear part of the competition curve for shellfish that contains between approximately 5 and 1500 µg STX equiv./100 g. Reagents were added in the following order: 35 µL STX standard or sample, then 35 µL [³H]STX, followed by 140 µL brain homogenate. The addition of brain homogenate was carried out with sufficient force to ensure mixing of the well contents, but without risk of splashing. The plate was then covered and incubated at 4°C for 1 h.

(b) *Assay filtration and counting.*—The plate was filtered using a microplate vacuum filtration manifold, and each well rinsed twice with 200 µL ice-cold HEPES buffer at a filtration rate that ensured all wells were dry within 2–5 s. The microplate was then placed in a microplate scintillation counter cassette, and the bottom was sealed with plate sealing tape. Lastly, 50 µL scintillation cocktail was added to each well, and the top of the plate was sealed with sealing tape. The plate was allowed to sit for 30 min to ensure impregnation of the filters with scintillant prior to counting for 1 min/well in the microplate scintillation counter.

Data Analysis

Curve fitting was performed using a four-parameter logistic curve fitting model for a one-site receptor binding using Wallac Multicalc software. The software reports the in-well sample concentration in nM equiv. STX. Sample concentration was then calculated in µg STX equivalents/100 g shellfish using the following formulas:

$$\begin{aligned}
 & (\text{nM equiv. STX}) \times (\text{sample dilution}) \times \frac{(210 \mu\text{L total volume})}{35 \mu\text{L sample}} \\
 & = \text{nM equiv. STX in extract}
 \end{aligned}$$

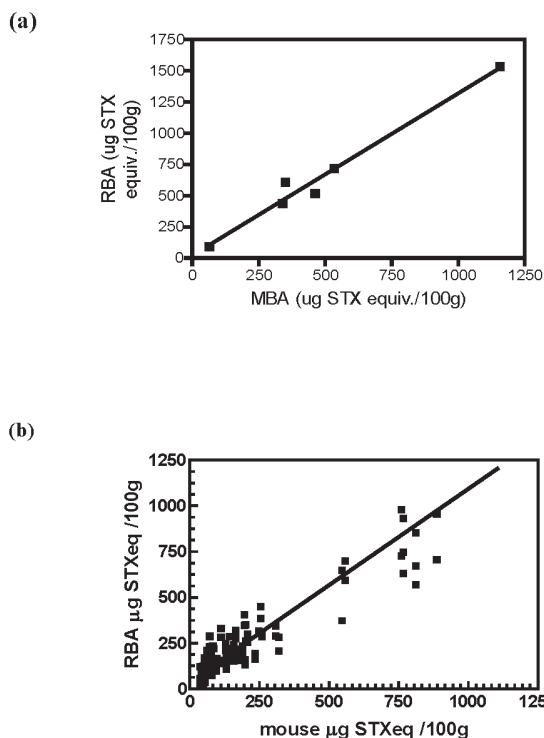


Figure 4. Linear correlation analysis between the RBA and mouse bioassay. (a) Average values of six naturally contaminated samples analyzed on five independent RBA assay days (r² = 0.98, slope = 1.29). (b) A separate study of 110 shellfish extracts analyzed by RBA and MBA yielded an r² of 0.88 with a slope of 1.32.

$$\begin{aligned}
 & (\text{nm equiv. STX in extract}) \times \frac{1 \text{ L}}{1000 \text{ mL}} \times \frac{372 \text{ ng}}{\text{nmol}} \times \frac{1 \mu\text{g}}{1000 \text{ ng}} \\
 & = \mu\text{g STX equiv./mL} \\
 & \mu\text{g STX equiv./mL} \times \frac{\text{mL extract}}{\text{g shellfish extracted}} \times 100 \\
 & = \mu\text{g STX equiv./100 g shellfish}
 \end{aligned}$$

Critical Control Points

(1) For a ligand that interacts specifically at one receptor site, the slope of the resulting competition curve should theoretically be 1.0. If the slope of the curve for a given assay is outside of the acceptable range of 0.8–1.2, linearity of the assay will be compromised, and quantification of the unknowns will be incorrect. Therefore, the assay should be re-run.

(2) The QC check standard should fall within ±30% of the stated value (3.0 nM). If the QC check standard does not fall within acceptable limits, the assay should be re-run.

Table 4. RBA-determined toxicities of nine naturally contaminated shellfish homogenates extracted using the 0.1 M HCl extraction method or the 1% acetic acid extraction method (μg STX equiv./100 g)

Sample	HCl			Acetic acid		
	Mean	SD	RSD	Mean	SD	RSD
1	11	4	36	19	7	39
2	600	143	24	488	104	21
3	690	142	21	584	167	29
4	136	8	6	131	41	31
5	152	27	18	167	21	13
6	302	87	29	270	72	27
7	340	88	26	264	63	24
8	262	79	30	252	48	19
9	63	26	41	54	19	34

(3) Sample quantification should be done only on dilutions that on the linear part of the curve [$b/b_0 = 0.2-0.7$, where B is the bound counts/min (CPM) in the sample and B_0 is the maximum CPM]. The RSD of the CPM must be $<30\%$.

(4) For a given sample, if none of the sample dilutions falls within the linear range (i.e., the concentration is too high, $b/b_0 < 0.2$), further dilutions must be made and the sample reanalyzed if a quantitative value is desired. If the sample concentration is too low to be quantified (i.e., $b/b_0 > 0.7$) at sample dilution 1:10, the sample must be reported as below the LOQ.

Mouse Bioassay and HPLC Procedures

Shellfish samples extracted in parallel using the HCl and acetic acid extraction methods described above were analyzed using the standard protocols prescribed by the AOAC methods for mouse bioassay (1) or precolumn oxidation HPLC method (2).

Results and Discussion

Calibration Curve

To establish the dynamic range and repeatability of the calibration curve, five assays were performed by one analyst on separate days. The composite curve (Figure 3) resulted in a half-maximal inhibition (IC_{50}) of $2.3 \text{ nM STX} \pm 0.3$ (RSD = 10.8%) with a slope of 0.96 ± 0.06 (RSD = 6.3%). Using the linear part of the curve ($0.2-0.7 b/b_0$) for quantification, a dynamic range of approximately one order of magnitude, $1.2-10.0 \text{ nM STX}$, was observed, as expected for a one-site binding assay. A QC check sample (3.0 nM STX) run in each assay averaged $3.0 \pm 0.5 \text{ nM}$ (RSD_r = 17.3%), with a recovery of 99.3% .

LOQ

Shellfish extracts were diluted a minimum of 10-fold prior to analysis to minimize matrix effects that can result in false positives. The LOQ was empirically determined as the

concentration, in a 10-fold diluted sample, that results in a b/b_0 of 0.7. This is a more conservative cutoff than the $0.8 b/b_0$ frequently used in receptor assays and was used because quantification was unacceptably variable above this b/b_0 cutoff. This results in an LOQ of approximately $5 \mu\text{g}$ equiv. STX/100 g shellfish, which provides a more than one order of magnitude margin relative to the regulatory limit of $80 \mu\text{g}/100 \text{ g}$.

Linearity

Linearity was assessed by five independent assays of three calibration standards that were expected to fall on the curve between 0.2 and $0.7 b/b_0$: $1.5, 3.0,$ and 6.0 nM STX prepared from FDA STX diHCl standard. Expected and measured values are listed in Table 1. Linear regression yielded a slope of 0.98 and an r^2 of 0.97 .

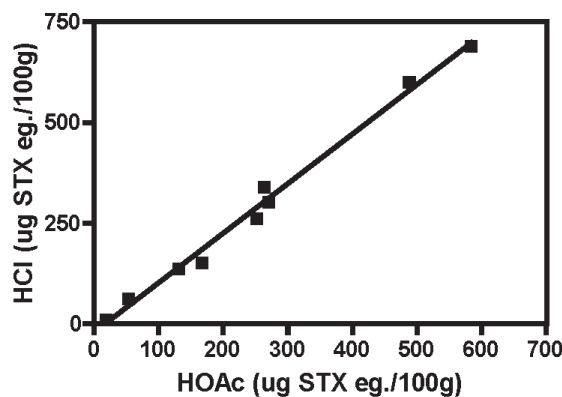


Figure 5. Linear correlation between HCl and acetic acid (HOAc) extracts analyzed by RBA. Results are average values of nine naturally contaminated samples obtained from four independent assays ($r^2 = 0.99$, slope = 1.23).

Table 5. HPLC analysis of nine naturally contaminated samples (1–9) extracted using 0.1 M HCl^a

Sample	STX	NEO ^b	GTX1,4 ^c	GTX2,3	B1	C1,2	Total PSP	As STX equivalent
HCl-1	3.5	0.0	0.0	7.3	0.0	0.0	10.8	6
HCl-2	231.6	23.9	42.6	324.5	28.3	249.7	900.4	412
HCl-3	220.8	53.7	74.9	436.3	43.4	338.1	1167.2	494
HCl-4	48.3	2.7	8.6	85.1	10.7	17.1	172.5	90
HCl-5	86.5	1.1	0.0	64.7	14.9	11.3	178.5	113
HCl-6	114.5	0.0	0.0	166.6	15.1	36.8	333.0	180
HCl-7	96.4	10.1	72.9	398.7	9.3	36.1	623.5	304
HCl-8	84.6	6.0	32.8	225.7	4.9	18.5	372.5	197
HCl-9	11.2	0.0	6.1	47.9	0.0	0.0	65.2	33

^a Values are in $\mu\text{g}/100\text{ g}$, as specific PSP congener or its STX equivalents, as indicated by the column headers.

^b NEO = Neosaxitoxin.

^c GTX = Gonyautoxin.

Recovery

Mussel tissue homogenates obtained from a local market were spiked with FDA STX diHCl standard at four levels bracketing the regulatory limit (0, 40, 80, and 120 $\mu\text{g}/100\text{ g}$) followed by thorough homogenization using a Polytron blender. Aliquots of spiked homogenate were stored at -80°C until extraction in 0.1 M HCl according to the protocol in the *Experimental* section. Extracts were analyzed in five assays performed on independent days. The mean recovery was 121% (Table 2).

Comparison of RBA-Reported Toxicity with the AOAC Mouse Bioassay

Six naturally contaminated shellfish samples were extracted in 0.1 M HCl according to the protocol in the *Experimental* section, and analyzed in five assays on

independent days (Table 3). Three shellfish species were represented: clam *Mya arenaria* (whole) LP1, LP4; mussel *Mytilus edulis* (whole) LP2, LP3; and scallop *Plactopecten magellanicus* (viscera) LP5, LP6. Between-assay RSDs ranged from 9 to 25% (mean 17.7%). An r^2 of 0.98 was obtained relative to the mouse bioassay, with a slope of 1.29 (Figure 4a).

A separate study of 110 naturally contaminated shellfish samples, extracted using the 0.1 M HCl method, and analyzed by RBA and mouse bioassay, yielded similar results with an r^2 of 0.88 and a slope of 1.32 (Figure 4b).

Effect of Extraction Method on RBA-Reported Toxicities

The recent approval of the precolumn oxidation HPLC method for PSP toxins as AOAC Official Method **2005.06** (3) and its potential recognition as a reference method for PSP

Table 6. HPLC analysis of the same nine naturally contaminated samples (1–9) extracted using 1% acetic acid^a

Sample	STX	NEO	GTX1,4	GTX2,3	B1	C1,2	Total PSP	As STX equivalent
HOAc-1	3.4	0.0	0.0	7.3	0.0	0.0	10.7	6
HOAc-2	187.6	13.1	21.7	280.7	25.1	248.9	777.1	329
HOAc-3	175.2	35.6	79.2	335.9	37.2	237.7	900.9	393
HOAc-4	33.4	3.1	11.3	61.8	6.0	15.5	131.1	68
HOAc-5	59.3	3.1	0.0	67.6	10.8	19.3	160.0	89
HOAc-6	100.8	0.0	0.0	158.0	11.8	28.4	299.0	162
HOAc-7	67.4	11.2	42.7	228.4	5.2	15.6	370.5	192
HOAc-8	71.0	8.3	34.4	190.3	4.3	12.6	320.8	173
HOAc-9	11.2	0.0	11.7	38.1	0.0	61.0	122.1	33

^a Values are in $\mu\text{g}/100\text{ g}$, as specific PSP congener or its STX equivalents, as indicated by the column headers.

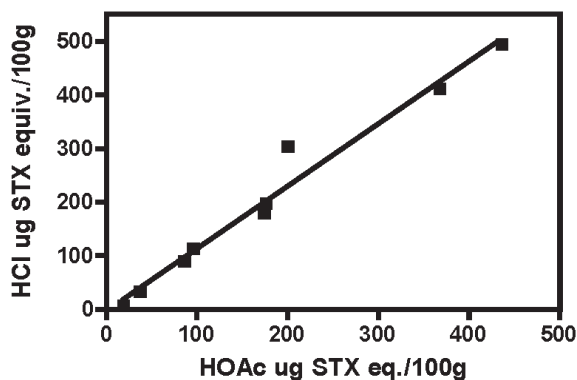
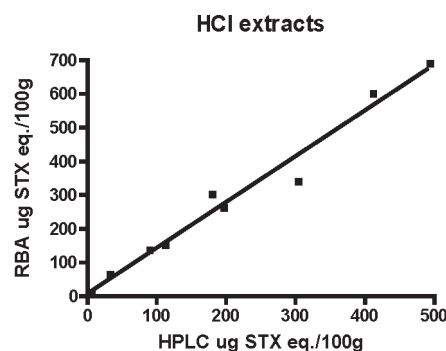


Figure 6. Linear correlation between HCl and acetic acid (HOAc) extracts analyzed by HPLC (slope = 1.16, $r^2 = 0.97$).

toxins prompted an investigation of the effects of extraction method on toxicity values reported by the RBA. Whereas the AOAC mouse bioassay prescribes shellfish extraction in 0.1 M HCl, the HPLC method uses extraction in 1% acetic acid. The 0.1 M HCl extraction procedure is known to result in the partial conversion of certain low-toxicity sulfocarbamoyl congeners to more highly toxic congeners in shellfish extracts, especially gonyautoxins, GTX5 and GTX6, to STX and neoSTX, and, thus, may result in somewhat higher toxicity values. To assess the effects of extraction procedure on RBA-reported toxicity, nine naturally contaminated shellfish samples (six blue mussel and three scallop) were homogenized and extracted independently using 0.1 M HCl and 1% acetic acid as described in the *Experimental* section. PSP toxicity in the extracts was then determined in four RBA assays run on independent days (Table 4). The between-assay RSD did not differ for samples prepared using the two extraction methods (25.8 and 26.3%, respectively). In general, the HCl extraction method resulted in slightly higher total toxicity values than reported for the acetic acid extracts (slope 1.23, $r^2 = 0.99$; Figure 5). The higher values reported for the HCl extracts are not explained by the conversion of sulfocarbamoyl toxins to more potent congeners in the HCl extracts, as can be seen in the toxin profiles determined by HPLC (Tables 5 and 6). Rather, the recovery of most congeners appears to be higher in the HCl extract. The higher concentrations reported in the HCl extract may reflect differences in the method by which volume is adjusted in the two extraction procedures. In the HCl method, final extract volume adjustment is made with the shellfish matrix present. In the acetic acid extraction, the matrix is first removed, the pellet re-extracted, the two extracts pooled, and then the final volume adjusted. HPLC analysis of the same samples showed a similar relationship between values reported for the HCl and acetic acid extracts (slope = 1.16, $r^2 = 0.97$; Figure 6) as seen in the RBA, with the HCl extracts containing greater STX equivalent/100 g.

(a)



(b)

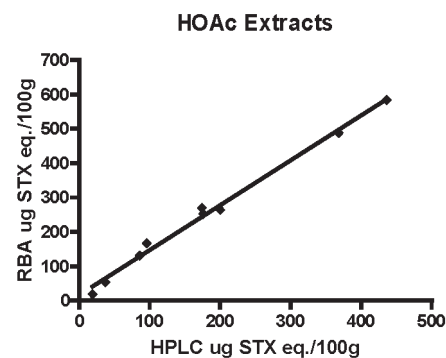


Figure 7. Linear correlation between RBA and HPLC for samples extracted (a) by the HCl method ($r^2 = 0.98$, slope = 1.39) and (b) by the acetic acid method ($r^2 = 0.99$, slope = 1.32).

Comparison of RBA with HPLC

The RBA showed good linear correlation with HPLC analysis of both HCl ($r^2 = 0.98$, slope = 1.39) and acetic acid ($r^2 = 0.99$, slope = 1.32) extracts, in both cases giving somewhat higher toxicities than the HPLC method (Figure 7). A number of factors may contribute to the difference in results for total toxic potencies by these two methods. The higher toxicity values given by the RBA may result in part from the fact that the HPLC method uses the STX free base molecular weight (300 Da), whereas the receptor assay (and mouse bioassay) uses the STX dihydrochloride molecular weight (372 Da) to calculate concentration, which would result in approximately 20% higher values in the RBA. Additional differences may result from the use of FDA as compared to the NRC saxitoxin standards in the RBA and HPLC methods, respectively. Higher RBA results may also result from the dominance of the more potent PSP congeners over the weaker congeners in mixtures competing for binding to the receptor, as detailed in ref. 13, which reflects their binding affinities. In

contrast to this complex behavior, the HPLC method adds linearly the concentrations of each congener based on toxic potencies determined by mouse bioassay for isolated congeners. In some cases, e.g., 11-hydroxysulfate epimers, the concentrations of separate epimers pairs are not resolved by HPLC, although their potencies differ widely as do their ratios in shellfish samples. Lastly, higher toxicity values reported by the RBA may reflect the presence of congeners or metabolites not reported by the HPLC method.

Ruggedness

Although formal ruggedness testing was not carried out during this SLV study, several steps in the procedure might be noted that can affect the precision and accuracy of the results. First, it is important to clarify shellfish extracts by centrifugation prior to running the assay, particularly if extracts are stored refrigerated or frozen before analysis, as precipitates in the extract may cause nonspecific binding that may result in overestimates of PSP toxin concentrations. Second, since the rat brain homogenate is a suspension, it is important to ensure that it remains evenly suspended by frequent vortex mixing or pipetting prior to and during its addition to the plate. The rate of assay plate filtration should ensure that the wells clear in 2–5 s, and the rinse buffer should be ice cold in order to minimize the rate of toxin release from the receptor. Lastly, following addition of liquid scintillant to the microplate wells, it is essential to allow a minimum of 30 min for the scintillant to penetrate the filters before counting. Counting prematurely can result in increased variability between wells and lower counts/well, thus increasing RSD. A count time of 1 min/well was chosen for this study as a compromise between optimum RSD and assay throughput. Increasing the count time to 5 min/well has been shown to improve the between-well RSD in this assay when using the Packard Top Count scintillation counter, a single detector instrument with somewhat lower efficiency than the Wallac Microbeta used in the current study (11).

Summary

This SLV and method comparison study demonstrates excellent linear correlation ($r^2 > 0.98$) between the microplate receptor binding assay and both the mouse bioassay and the precolumn oxidation HPLC method for the determination of PSP toxins in shellfish. The microplate format of the assay, when coupled with microplate scintillation counting, provides a quantitative high throughput screening tool for PSP toxin testing in shellfish. The tendency of the RBA to overestimate PSP toxicity relative to the reference methods minimizes the chance of returning false negatives. Where RBA-measured

toxicity results in STX equivalent values close to the regulatory limit, confirmation with a reference method is necessary if a regulatory decision is being made. Nonetheless, application of the assay as a high throughput screen can alleviate the unnecessarily large numbers of animals used for the mouse bioassay on negative samples and, similarly, alleviate the lengthy analysis of samples by HPLC at very high or very low concentrations. We propose that this method be collaboratively tested to establish if it is robust enough to be used in monitoring and regulatory laboratories.

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FOOD CHEMICAL CONTAMINANTS**Determination of Paralytic Shellfish Toxins in Shellfish by Receptor Binding Assay: Collaborative Study**

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A collaborative study was conducted on a microplate format receptor binding assay (RBA) for paralytic shellfish toxins (PST). The assay quantifies the composite PST toxicity in shellfish samples based on the ability of sample extracts to compete with ^3H saxitoxin (STX) diHCl for binding to voltage-gated sodium channels in a rat brain membrane preparation. Quantification of binding can be carried out using either a microplate or traditional scintillation counter; both end points were included in this study. Nine laboratories from six countries completed the study. One laboratory analyzed the samples using the precolumn oxidation HPLC method (AOAC Method 2005.06) to determine the STX congener composition. Three laboratories performed the mouse bioassay (AOAC Method 959.08). The study focused on the ability of the assay to measure the PST toxicity of samples below, near, or slightly above the regulatory limit of 800 (μg STX diHCl equiv./kg). A total of 21 shellfish homogenates were extracted in 0.1 M HCl, and the extracts were analyzed by RBA in three assays on separate days. Samples included naturally contaminated shellfish samples of different species collected from several geographic regions, which contained varying STX congener profiles due to their exposure to different PST-producing dinoflagellate species or differences in toxin metabolism: blue mussel (*Mytilus edulis*) from the U.S. east and west coasts, California mussel (*Mytilus californianus*) from the U.S. west coast, chorito mussel (*Mytilus chilensis*) from Chile, green mussel (*Perna canaliculus*) from New Zealand,

Atlantic surf clam (*Spisula solidissima*) from the U.S. east coast, butter clam (*Saxidomus gigantea*) from the west coast of the United States, almeja clam (*Venus antiqua*) from Chile, and Atlantic sea scallop (*Plactopecten magellanicus*) from the U.S. east coast. All samples were provided as whole animal homogenates, except Atlantic sea scallop and green mussel, from which only the hepatopancreas was homogenized. Among the naturally contaminated samples, five were blind duplicates used for calculation of RSD_r . The interlaboratory RSD_R of the assay for 21 samples tested in nine laboratories was 33.1%, yielding a HorRat value of 2.0. Removal of results for one laboratory that reported systematically low values resulted in an average RSD_R of 28.7% and average HorRat value of 1.8. Intralaboratory RSD_r , based on five blind duplicate samples tested in separate assays, was 25.1%. RSD_r obtained by individual laboratories ranged from 11.8 to 34.9%. Laboratories that are routine users of the assay performed better than nonroutine users, with an average RSD_r of 17.1%. Recovery of STX from spiked shellfish homogenates was 88.1–93.3%. Correlation with the mouse bioassay yielded a slope of 1.64 and correlation coefficient (r^2) of 0.84, while correlation with the precolumn oxidation HPLC method yielded a slope of 1.20 and an r^2 of 0.92. When samples were sorted according to increasing toxin concentration (μg STX diHCl equiv./kg) as assessed by the mouse bioassay, the RBA returned no false negatives relative to the 800 μg STX diHCl equiv./kg regulatory limit for shellfish. Currently, no validated methods other than the mouse bioassay directly measure a composite toxic potency for PST in shellfish. The results of this interlaboratory study demonstrate that the RBA is suitable for the routine determination of PST in shellfish in appropriately equipped laboratories.

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Paralytic shellfish poisoning (PSP) is caused by a suite of heterocyclic guanidinium toxins collectively called saxitoxins (STXs). Currently more than 21 congeners of STX are known; they occur in varying proportions in the dinoflagellates that produce them and may be further

Table 1. Shellfish homogenate samples analyzed for PSTs in the collaborative study^a

Sample No.	Sample ID	Shellfish species and origin	Blind duplicate
1	MLV05	Atlantic sea scallop (<i>Plactopecten magellanicus</i>) from the U.S. east coast	x
2	MLV06	California mussel (<i>Mytilus californianus</i>) from the U.S. west coast	x
3	MLV08	Green mussel (<i>Perna canaliculus</i>) from New Zealand	
4	MLV09	Blue mussel (<i>M. edulis</i>) from the U.S. west coast	x
5	MLV12	Blue mussel (<i>M. edulis</i>) east coast U.S., spiked with 200 µg/kg STX diHCl	
6	MLV14	Blue mussel (<i>M. edulis</i>) east coast U.S., spiked with 1200 µg/kg STX diHCl	
7	MLV16	Almeja clam (<i>Venus antique</i>) from Chile	
8	MLV01	Surf clam (<i>Spisula solidissima</i>) from the U.S. east coast	
9	MLV02	Chorito mussel (<i>M. chilensis</i>) from Chile	
10	MLV04	Scallop (<i>Plactopecten magellanicus</i>) from the U.S. east coast	
11	MLV07	Blue mussel (<i>M. edulis</i>) east coast U.S.	x
12	MLV09	Blue mussel (<i>M. edulis</i>) from the U.S. west coast	x
13	MLV11	Almeja clam (<i>Venus antique</i>) from Chile clam	x
14	MLV13	Blue mussel (<i>M. edulis</i>) east coast U.S., spiked with 500 µg/kg STX diHCl	
15	MLV03	Chorito mussel (<i>M. chilensis</i>) from Chile	
16	MLV05	Atlantic sea scallop (<i>Plactopecten magellanicus</i>) from the U.S. east coast	x
17	MLV06	California mussel (<i>M. californianus</i>) from the U.S. west coast	x
18	MLV07	Blue mussel (<i>M. edulis</i>) east coast U.S.	x
19	MLV10	Butterclam (<i>Saxidomus gigantea</i>) from the U.S. west coast	
20	MLV11	Almeja clam (<i>Venus antique</i>) from Chile clam	x
21	MLV15	Blue mussel (<i>M. edulis</i>) negative control, east coast U.S.	

^a Sample number identifies the individual samples analyzed in the assays, with 1–7 analyzed in assay 1, 8–14 in assay 2, and 15–21 in assay 3. Sample identification (MLV for multilaboratory validation) describes the 16 unique samples, among which five were assayed as blind duplicates, to make a total of 21 samples. Blind duplicates, run in different assays, are identified by an “x.”

metabolized in shellfish that accumulate them, making analytical determination of paralytic shellfish toxins (PST) in shellfish complex. The long-standing regulatory method for PST is the AOAC mouse bioassay (1; AOAC Method **959.08**), with a regulatory limit of 800 µg STX di HCl equiv./kg shellfish generally applied, but established at 400 µg STX diHCl equiv./kg in certain countries (e.g., the Philippines). However, at concentrations near the regulatory limit, the mouse bioassay can significantly underestimate PST in shellfish (2). This, in addition to increasing resistance to live animal testing in both the United States and the European Union (EU), has increased the need to develop alternative methods suitable for use in a high-throughput monitoring or regulatory setting.

In the past decade, several alternatives to the mouse bioassay have been developed. In the EU, the mouse bioassay remains the reference method for PST in shellfish, but European Commission (EC) Regulation 1664/2006 specifies that other internationally recognized methods may be used. Two HPLC methods, a precolumn oxidation method (3, 4; AOAC Method **2005.06**) and a postcolumn oxidation method (5; AOAC Method **2011.02**), have been approved by AOAC as *Official Methods*SM for PSP toxin analysis. The EC directive recognizes the precolumn oxidation HPLC method (AOAC Method **2005.06**) as an alternative to the mouse bioassay, but retains the mouse bioassay as the reference method in instances where results are challenged. HPLC methods separate and quantify individual

STX congeners, which are then recombined according to their toxic equivalencies to yield a composite PST toxicity value. Although the HPLC methods perform well quantitatively, a high-throughput screening method capable of reporting toxic potency directly is still desirable for monitoring programs that often screen large numbers of negative samples. A qualitative lateral flow antibody test for PST with a reported detection limit of 400 µg STX equiv./kg was developed by Jellett Rapid Testing Ltd (Chester Basin, NS, Canada) and approved by the U.S. Interstate Shellfish Sanitation Conference and the U.S. Food and Drug Administration as a screening method in specific circumstances. This method performed well in a comparison study with the mouse bioassay (6), but is not fully quantitative and has not been subjected to a full AOAC collaborative trial. To date, a suitable quantitative, high-throughput alternative to the mouse bioassay has not been validated through the AOAC Official Methods Program. The receptor binding assay (RBA) for PST is an excellent candidate for fulfilling the requirements of a high-throughput, quantitative assay that directly reports a composite toxic potency.

The basis of the RBA is the interaction between the toxins and their pharmacological target. All STX congeners bind to site 1 on the alpha subunit of the voltage-gated sodium channel with binding affinities proportional to their toxic potency (7). Therefore, an RBA can quantitatively measure the combined toxic potency of mixtures of STX congeners in a sample,

independent of the toxin congeners present (8). In the RBA for PST, tritiated STX ($[^3\text{H}]$ STX) competes with unlabeled STX and/or its congeners for a finite number of available receptor sites in a rat brain membrane preparation. Following establishment of binding equilibrium, unbound $[^3\text{H}]$ STX is removed by filtration and receptor bound $[^3\text{H}]$ STX quantified by liquid scintillation counting. The reduction in $[^3\text{H}]$ STX binding is directly proportional to the amount of unlabeled toxin present. A standard curve is generated using increasing concentrations of nonradiolabeled STX standard from 10^{-10} to 10^{-6} M STX. The concentration of toxin in samples is determined in reference to the standard curve.

The assay being tested in this collaborative trial is a modification of the method of Doucette et al. (9) to incorporate a 96-well microtiter plate format, which increases sample throughput and minimizes error by reducing sample handling and pipetting steps. This microplate PST RBA was evaluated in a single-laboratory validation (SLV) study (10), which established an interassay repeatability (RSD_r) of 17.7% and good correlation with the mouse bioassay and precolumn oxidation HPLC methods. The toxin concentrations in shellfish tested in the SLV study ranged from near to well above the regulatory limit (approximately 900–15 000 μg STX diHCl equiv./kg). The current study focuses more specifically on the performance of the RBA in the critical range of shellfish toxicities below, near, and slightly above the regulatory limit (approximately 150–2400 μg STX diHCl equiv./kg).

The results of the collaborative study suggest that the RBA for PST is a suitable high-throughput screen for PST in shellfish. Although HPLC methods offer quantitative information on congener composition of samples, often the desired information is composite toxic potency, which requires the summation of individual congeners, corrected for their individual toxic equivalencies. The RBA provides a single integrated toxic potency value that reflects activity of all known and potential unknown congeners present in the sample. Use of the microtiter plate format, in conjunction with microplate scintillation counting, provides the ability to screen multiple samples simultaneously in a total assay time of less than 3 h. The assay format described in the current study provides for the quantitative determination of composite PST toxicity in seven shellfish extracts per 96-well microplate, each run in triplicate at three dilutions, covering toxicity ranges of approximately 35–6000 μg STX diHCl equiv./kg. In a high-throughput assay setting, multiple plates can be set up simultaneously, so that six assay plates can easily be accommodated each day by a single analyst, for a throughput of 42 samples/day. This compares favorably to an estimated throughput of 20–25 samples a day by the precolumn HPLC method (B. Niedzwiadek, Health Canada, personal communication) or 30–35 by mouse bioassay (B. Suarez, University of Chile, personal communication).

Collaborative Study

The focus of this study was to assess the performance of the RBA to determine PST toxicity in samples of commercially important shellfish at a range of concentrations below and above the regulatory limit. Twenty-one shellfish homogenates were included in the study, which represented 16 unique samples (Table 1). The homogenates included 12 naturally contaminated shellfish samples of different species collected from several

geographic regions: blue mussel (*M. edulis*) from the U.S. east and west coasts, California mussel (*M. californianus*) from the U.S. west coast, chorito mussel (*M. chilensis*) from Chile, green mussel (*Perna canaliculus*) from New Zealand, Atlantic surf clam (*Spisula solidissima*) from the U.S. east coast, butter clam (*Saxidomus gigantea*) from U.S. west coast, almeja clam (*Venus antiqua*) from Chile, and Atlantic sea scallop (*Plactopecten magellanicus*) from the U.S. east coast. All samples were provided as whole animal homogenates, except Atlantic sea scallop and green mussel, which included hepatopancreas only. Among the naturally contaminated samples, five were blind duplicates tested on separate days that were used for calculation of RSD_r . Samples run as duplicates are indicated in Table 1. Three samples consisting of STX-spiked mussel homogenate (*M. edulis*) at levels that bracketed the regulatory limits of 800 $\mu\text{g}/\text{kg}$ (500 and 1200 $\mu\text{g}/\text{kg}$ spike) and 400 $\mu\text{g}/\text{kg}$ (200 $\mu\text{g}/\text{kg}$ spike) were included to calculate recovery. One sample was the negative control homogenate of *M. edulis* to which the STX spikes were added. All homogenates were extracted by the study participants and the extracts analyzed by RBA in three assays on separate days.

Study Participants

Ten laboratories from seven countries agreed to carry out RBAs for this study, including the United States, Italy, Australia, New Zealand, Thailand, the Philippines, and South Africa. Participants included laboratories from regulatory authorities, as well as government and academic laboratories with monitoring needs. Five of the participating laboratories (Laboratories 1–5) have this method well established and may be considered routine users. Two laboratories had previous experience running this format of the PST RBA, but have not implemented it routinely. One laboratory had previous experience with receptor assays, but had not used the microplate filtration format of the assay. One laboratory had no previous experience with RBAs. Three laboratories from different countries, United States, Chile, and Thailand, carried out the AOAC official mouse bioassay method (AOAC Method 959.08) on the same set of samples. All mouse bioassay laboratories were experienced regulatory authorities with monitoring responsibilities. One laboratory (Health Canada) performed the precolumn oxidation HPLC method for PST (AOAC Method 2005.06).

Preparation of Homogenates

All shellfish samples were thoroughly homogenized using a polytron blender. For spiked samples, saxitoxin standard reference material (STX diHCl) was added to the specified concentration, and the sample was thoroughly rehomogenized to ensure homogeneity. The toxin congener profiles and concentrations of all samples were determined by the precolumn oxidation HPLC method (performed by Health Canada). STX equivalents were determined by mouse bioassay (performed by Maine Department of Marine Resources). Subsamples of each homogenate (12 g) were packaged in polycarbonate tubes and stored at -80°C until shipment to collaborating laboratories by courier. All samples were coded prior to distributing to collaborating laboratories, with the codes to each laboratory being unique, and provided blind. Coding consisted of two letters followed by a number in the form X A1-7, X B1-7, and

X C1-7, where the X indicated the laboratory, the second letter indicated the three assays to be conducted, and the numerical code indicated sample number within that assay. Three practice homogenates were similarly produced.

Shipment of Study Material

The following reagents were provided to the collaborating laboratories in a single shipment containing enough dry ice to keep the contents frozen for 5 days: [³H] STX; STX diHCl standard; rat brain membrane preparation; 21 coded shellfish homogenates; three practice homogenates; and a QC check sample consisting of 18 nM STX diHCl. Sufficient homogenate (12 g) was provided to ensure an accurate weight of material could be removed from the storage vial if an additional extraction were necessary due to unexpected circumstances. The identity of the samples was not released to collaborators. All reagents were received frozen and in good condition. Each participant received electronically a detailed assay protocol, comprehensive instructions for conducting the study and data reporting, and data reporting forms.

Analysis

Participants extracted all homogenates using a modification of the 0.1 M HCl extraction method used in the AOAC standard mouse bioassay protocol (modified only by scale). They were asked to perform three RBAs, each on separate days. Each assay consisted of one 96-well plate that included a standard curve, QC check sample, and seven shellfish extracts. All samples and standards were tested in triplicate wells. All shellfish extracts were run at three dilutions (1/10, 1/50, and 1/200), which ensured that at least one dilution would fall on the linear part of the standard curve. Participants were instructed to analyze samples coded A, B, or C in the first, second, or third assay, respectively, in numerical order. The five blind duplicate samples were coded so that they were tested in two independent assays, with the combination of assays differing between duplicates. Before performing the official study, participants were asked to run a practice assay that included three shellfish homogenates in the same format to ensure that any unexpected problems were encountered and addressed prior to the official study. The practice samples consisted of a negative control mussel homogenate (MLV15), and two naturally contaminated samples that were also included in the full study (MLV05 and MLV11). The identity of the practice samples was not made known to participants. Results of the practice run were submitted by e-mail to the coordinating laboratory for review before proceeding with the full study.

For the mouse bioassay, participants followed the AOAC official mouse bioassay method (AOAC Method 959.08), with the exception of a modified 0.1 M HCl extraction protocol used in the RBA protocol, which was modified only by scale so that 5 mL 0.1 M HCl was added to 5 g of shellfish homogenate, with all other aspects of the extraction protocol being identical. The HPLC laboratory followed the precolumn oxidation HPLC method for PST (AOAC Method 2005.06); however, final concentrations in µg/kg and µg STX equiv./kg were calculated using the formula weight of STX diHCl [372 daltons (da)], as opposed to the free base (299.3 da) in the standard HPLC protocol, to more directly compare with the RBA.

Data Analysis and Reporting

Participants were asked to report whether they used a standard or microplate scintillation counter for the study and, if a microplate counter was used, which model, because of differences in inherent counting efficiency between current commercially available counters. For data analysis, participants were instructed to use GraphPad Prism software (La Jolla, CA) or the on-board curve-fitting software provided with their microplate scintillation counter e.g., PerkinElmer Wallac MultiCalc (Gaithersburg, MD) or Packard Top Count software (Packard Instrument Co., Meriden, CT), and to report what software was used. For analysis, a four parameter logistic fit, also known as a sigmoidal dose response with variable slope, or Hill equation, was prescribed. Participants presented their analyzed data on the spreadsheet template provided, including assay quality parameters (slope, IC₅₀, and quantification of the QC check sample), between-well CVs for each sample dilution that fell within the linear part of the standard curve (0.2–0.7 B/B₀), and calculated values for these samples in the well (nM), in the extract (µg STX equiv./mL), and in the shellfish tissue (µg STX equiv./kg). Participants were also asked to report all raw count data so that all results could be analyzed by the coordinating laboratory using identical software (GraphPad Prism 4.0) to assess whether systematic differences in quantification arose from using different curve-fitting software. All data were reported via e-mail to the coordinating laboratory.

The calculated results sheets were reviewed by the coordinating laboratory for obvious errors in dilutions and calculations and for use of the prescribed curve-fitting model. Obvious errors were corrected and the participant laboratory was consulted for concurrence. The reviewed results were then used for evaluation in the collaborative study.

Statistical Evaluation of the Collaborative Study

For each sample analyzed, outliers were first determined using the Grubbs test at a probability value of 1% (www.graphpad.com), with no more than one outlier removed, so that valid data remained from a minimum of eight laboratories. The mean, S_R, and RSD_R, and HorRat values were then calculated for each sample. For blind duplicates, the AOAC INTERNATIONAL Interlaboratory Study Workbook for Blind Duplicates, v2.0, was used to further evaluate for outliers and determine S_r and RSD_r. GraphPad Prism was used to determine correlation among the RBA, mouse bioassay, and HPLC results.

AOAC Official Method 2011.27 Paralytic Shellfish Toxins (PSTs) in Shellfish Receptor Binding Assay First Action 2011

[Applicable to the determination of paralytic shellfish toxins (PSTs), as µg STX diHCl equiv./kg, in shellfish (mussels, clams, scallops) at levels >149 µg STX diHCl equiv./kg, with a limit of detection (LOD) of 45 STX diHCl equiv./kg shellfish and a limit of quantification (LOQ) of 126 µg STX diHCl equiv./kg shellfish.]

Caution: Wear disposable gloves and protective laboratory coat while performing the assay. PSTs are neurotoxins that are harmful if ingested. The assay uses a tritium labeled tracer, [³H] STX, at low concentration.

All laboratories performing the assay must have approved radiation laboratory space and must follow procedures prescribed by their nuclear regulatory agency for receipt, use, and disposal of isotopes.

See Tables 2011.27A–E for results of the interlaboratory study supporting acceptance of the method.

A. Principle

Test portions of shellfish homogenates are extracted using the AOAC mouse bioassay extraction protocol (959.08), modified by scale. The PST receptor assay is a competitive binding assay in which [³H] STX competes with unlabeled STX in standards or mixtures of PST in samples for a finite number of available receptor sites (site 1 on the voltage gated sodium channel) in a rat brain membrane preparation. Following establishment of binding equilibrium at 4°C, unbound [³H] STX is removed by filtration and bound [³H] STX is quantified by liquid scintillation counting. A standard curve is generated using increasing concentrations of STX standard from 10⁻¹⁰ to 10⁻⁶ M STX, which results in a reduction in bound [³H] STX that is directly proportional to the amount of unlabeled toxin present. The concentration of toxin in samples is determined in reference to the standard curve. Incubation is carried out in a microplate format to minimize sample handling and the amount of radioactivity used. Bound [³H] STX (as counts per minute; CPM) can be determined either by conventional or by microplate scintillation counting. Both methods are included in this protocol.

B. Apparatus and Supplies

- (a) *Traditional or microplate scintillation counter.*
 - (b) *Micropipettors.*—1–1000 µL variable volumes and disposable tips.
 - (c) *Eight channel pipettor.*—5–200 µL variable volume and disposable tips.
 - (d) *96-Well microtiter filter plate.*—With 1.0 µm pore size type GF/B glass fiber filter/0.65 µm pore size Durapore support membrane (Millipore, Bedford, MA; Cat. No. MSFB N6B 50).
 - (e) *MultiScreen vacuum manifold.*—Millipore; Cat. No. NSVMHTS00.
 - (f) *Vacuum pump.*
 - (g) *Centrifuge tubes.*—15 and 50 mL, conical, plastic.
 - (h) *Mini dilution tubes in 96-tube array.*
 - (i) *Reagent reservoirs.*
 - (j) *Ice bucket and ice.*
 - (k) *Vortex mixer.*
 - (l) *Sealing tape.*—Millipore; Cat. No. MATA HCL00.
 - (m) *Volumetric flask.*—1 L.
 - (n) *-80°C freezer.*
 - (o) *Refrigerator.*
- For traditional scintillation counter only:
- (p) *MultiScreen punch device.*—Millipore; Cat No. MAMP 096 08.
 - (q) *MultiScreen disposable punch tips.*—Millipore; Cat. No. MADP 196 10.
 - (r) *MultiScreen punch kit B for 4 mL vials.*—Millipore; Cat. No. MAPK 896 0B.
 - (s) *Scintillation vials.*—4 mL.
- For sample extraction:
- (t) *Pipets.*
 - (u) *Centrifuge tubes.*—15 mL, conical, plastic.

- (v) *Vacuum pump or house vacuum.*
- (w) *pH meter or pH paper.*
- (x) *Hot plate.*
- (y) *Graduated centrifuge tubes.*—15 mL.
- (z) *Centrifuge and rotor for 15 mL tubes.*

C. Reagents

- (a) [³H] STX.—0.1 mCi/mL, ≥10 Ci/mmol, ≥90% radiochemical purity (American Radiolabeled Chemicals, St. Louis, MO, or International Isotopes Clearinghouse, Leawood, KS).
 - (b) *STX diHCl.*—NIST RM 8642 (www.nist.gov).
 - (c) *3-Morpholinopropanesulfonic acid (MOPS).*—Sigma (St. Louis, MO; Cat. No. M3183-500G), or equivalent.
 - (d) *Choline chloride.*—Sigma (Cat. No. C7527-500G), or equivalent.
 - (e) *Rat brain membrane preparation.*—See Appendix.
- For traditional counter:
- (f) *Scintiverse BD liquid scintillation cocktail.*—Fisher Scientific (Waltham, MA; Cat. No. SX-18), or equivalent.
- For microplate counter:
- (g) *Optiphase liquid scintillation cocktail.*—PerkinElmer Life Sciences (Downers Grove, IL; Cat. No. 1200-139), or equivalent.
- For sample extraction:
- (h) *Hydrochloric acid (HCl).*—1.0 and 0.1 M.
 - (i) *Sodium hydroxide.*—0.1 M.
 - (j) *Water.*—Distilled or deionized (18 µΩ).

D. Sample Extraction

Accurately weigh 5.0 g tissue homogenate into a tared 15 mL conical tube. Add 5.0 mL of 0.1 M HCl, vortex, and check pH. If necessary, adjust pH to 3.0–4.0 as determined by a pH meter or pH paper. To lower pH, add 1 M HCl dropwise with mixing; to raise pH, add 0.1 M NaOH dropwise with mixing to prevent local alkalization and consequent destruction of toxin. Place the tube in a beaker of boiling water on a hot plate for 5 min with the caps loosened. Remove and cool to room temperature. Check pH and adjust cooled mixture to pH 3.0–4.0 as described above. Transfer entire contents to graduated centrifuge tube and dilute volumetrically to 10 mL. Gently stir contents to homogeneity and allow to settle until portion of supernatant is translucent and can be decanted free of solid particles. Pour approximately 5 to 7 mL of the translucent supernatant into a centrifuge tube. Centrifuge at 3000 × g for 10 min. Retain clarified supernatant and transfer to a clean centrifuge tube. Store extracts at -20°C until tested in receptor assay.

E. Preparation of Stock Solutions and Standards

- (a) *Assay buffer.*—100 mM MOPS/100 mM choline chloride, pH 7.4. Weigh out 20.9 g MOPS and 13.96 g choline chloride and add to 900 mL dH₂O. Adjust pH to 7.4 with NaOH while stirring and bring to a final volume of 1 L with dH₂O. Store at 4°C.
- (b) *Radioligand solution.*—Calculate the concentration of [³H] STX stock provided by the supplier, which may vary between lots. Suppliers generally provide the specific activity in Ci/mmol (generally 10–30 Ci/mmol) and activity in mCi/mL (0.05–0.1 mCi/mL), from which the molar concentration can be calculated. Prepare 4 mL of a 15 nM working stock of [³H] STX fresh daily in 100 mM MOPS/100 mM choline chloride

Table 2011.27A. Receptor binding assay results on individual samples (values are in µg STX diHCl equiv./kg shellfish tissue); summary statistics on all samples; summary statistics excluding Laboratory 9

Assay	No.	Sample															All labs				Labs 1-8		
		Lab															Mean	S _R	RSD _R , %	HorRat	Mean	S _R	RSD _R , %
		1	2	3	4	5	6	7	8	9	Mean	S _R	RSD _R , %	HorRat	Mean	S _R	RSD _R , %	HorRat					
Day 1	1	MLV05	370	610	620	410	690	1070	630	660	330	599	222	37.1	2.2	633	212	33.5	2.0				
	2	MLV06	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	422	28.6	1.9	1549	386	24.9	1.7				
	3	MLV08	80	190	140	90	130	160	230	220	100	149	55	37.2	1.8	155	56	36.0	1.7				
	4	MLV09	860	680	950	870	980	1120	1460	820	590	926	255	27.5	1.7	968	237	24.5	1.5				
	5	MLV12	180 ^a	200	200	150	150	100	290	290	100	168	62	37.2	1.8	177	60	34.1	1.7				
	6	MLV14	950	940	1060	1130	1040	750	1460	1320	810	1051	228	21.7	1.4	1081	224	20.7	1.3				
	7	MLV16	660	930	1080	870	840	1320	1490	2420 ^b	490	960	329	34.3	2.1	1027	291	28.3	1.8				
Day 2	8	MLV01	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	818	48.6	3.3	1829	739	40.4	2.8				
	9	MVL02	830	1180	1130	1150	1130	1780	1340	980	690	1134	311	27.4	1.8	1190	281	23.6	1.5				
	10	MLV04	2440	2840	2910	1740	2150	1810	2690	2490	1210	2253	572	25.4	1.8	2384	446	18.7	1.3				
	11	MLV07	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	345	23.9	1.6	1520	279	18.3	1.2				
	12	MLV09	810	1190	1130	810	1630	1390	1880	1120	870	1203	372	30.9	2.0	1245	375	30.1	2.0				
	13	MLV11	270	370	480	340	640	490	240	600	110	393	174	44.3	2.4	429	148	34.4	1.9				
	14	MLV13	400	1240 ^b	560	450	650	530	500	440	200	466	133	28.5	1.6	504	85	16.8	1.0				
Day 3	15	MLV03	330	270	410	180	590	680	370	1570 ^b	90	365	197	54.0	2.9	404	176	43.5	2.4				
	16	MLV05	580	670	250	430	910	700	860	940	300	627	257	41.1	2.4	668	242	36.2	2.1				
	17	MLV06	1290	1520	1460	970	1800	2520	1470	870	1250	1461	488	33.4	2.2	1488	515	34.6	2.3				
	18	MLV07	1010	1600	1390	1000	1720	1860	1520	2150	890	1460	429	29.4	2.0	1531	397	26.0	1.7				
	19	MLV10	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	570	24.0	1.7	2443	569	23.3	1.7				
	20	MLV11	430	350	460	280	550	620	1149 ^b	410	250	419	127	30.2	1.7	443	115	26.0	1.4				
	21	MLV15	ND ^c	ND	ND	ND	ND	ND	ND	180	ND	—	—	—	—	—	—	—	—				
	Avg. RSD _R													33.2					28.7				
	Avg. HorRat													2.0					1.8				

^a CV 41%; not used in calculations.

^b Outlier; not used in calculations.

^c ND = Not detected.

Table 2011.27B. Summary statistics on blind duplicates, run in separate assays (values are in μg STX diHCl equiv./kg)

Lab	MLV05		MLV06		MLV07		MLV09		MLV11		Avg.
	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	Assay 1	Assay 2	
1	370	580	1100	1290	1260	1010	860	810	270	430	
2	610	670	1340	1520	1540	1530	680	1190	370	350	
3	620	250	1320	1460	1220	1390	950	1130	480	401	
4	410	430	1440	970	1980	1000	870	810	340	280	
5	690	910	1260	1790	1760	1720	980	1630	640	550	
6	1070	700	1720	2520	1530	1860	1120	1390	490	620	
7	630	880	2090	1240	1750	1150	1460	1830	230 ^a	1149 ^a	
8	660	940	2130	870	1210	2150	820	1120	600	410	
9	330	300	890	1250	840	890	590	870	110	250	
Avg.		614		1453		1433		1062		416	
S _r		169		432		366		247		83	
S _R		239		444		387		338		152	
RSD _r , %		27.5		29.4		25.5		23.3		20.0	25.1
RSD _R , %		38.9		30.2		27.0		31.9		36.5	32.9
HorRat		2.3		2.0		1.8		2.0		2.0	2.0

^a Outlier; not used in calculation.

buffer. This will provide sufficient volume for one 96-well plate at an in-well concentration of 2.5 nM. Measure total counts of each working stock prior to running an assay: add 35 μL of the working stock [^3H] STX in buffer to a liquid scintillation vial with 4 mL scintillant and count on a traditional liquid scintillation counter. This is done to confirm correct dilution prior to running the assay. Depending on the efficiency of the scintillation counter used, the corresponding CPM will vary, but should be consistent day-to-day and within 15% of the expected value.

(c) *Unlabeled STX standard working solution.*—The STX diHCl standard is provided at a concentration of 268.8 μM (100 $\mu\text{g}/\text{mL}$). A “bulk” standard curve can be made up in advance and stored at 4°C for up to 1 month. The use of a bulk standard curve minimizes the pipetting needed for setting up an assay routinely and improves day-to-day repeatability. Make up 3 mM HCl (e.g., from a 3 M stock, 50 μL in 50 mL), then perform the serial dilutions (see Table 2011.27F) of NIST RM 8642 STX diHCl (100 $\mu\text{g}/\text{mL}$ = 268.8 μM) to make up the standard curve in 3 mM HCl. These standard stock solutions will be diluted 1/6 in the assay to yield the designated in-assay concentrations (see Table 2011.27F).

(d) *Interassay calibration standard (QC check).*—Prepare a reference standard containing 1.8×10^{-8} M STX standard (3.0×10^{-9} M STX in assay) in advance in 3 mM HCl and keep frozen (−80°C) in 1 mL aliquots for long-term storage. Aliquots should be thawed and stored at 4°C for routine use (stable up to 1 month) and analyzed in each assay. This serves as a QC check and confirms day-to-day performance of the assay.

(e) *Rat brain membrane preparation.*—Prepare rat brain membrane preparation in bulk (see Appendix: *Rat Brain Membrane Preparation*) and store at −80°C until used in the assay. Thaw an aliquot of rat brain membrane preparation on ice. Dilute membrane preparation with cold (4°C) 100 mM

MOPS/100 mM choline chloride, pH 7.4, to yield a working stock with a protein concentration of 1.0 mg/mL (this will be diluted in the assay plate to 0.5 mg/mL in-well concentration). Vortex vigorously to achieve a visibly homogeneous suspension. Keep the diluted membrane preparation on ice until ready to use.

F. Performing the Assay

(a) *Plate setup.*—When possible, use a multichannel pipet to minimize pipetting effort and increase consistency. Standard curve, QC check, and sample extracts are run in triplicate wells. Multiple dilutions of sample extracts should be analyzed in order to obtain a value that falls between 0.2–0.7 B/B₀ on the standard curve for quantification. For ease of analysis, it is convenient to use a standard plate layout that maximizes the number of samples and standards that can be analyzed on one plate. For shellfish extracts, a minimum dilution of 1:10 is used, which minimizes potential matrix effects, while still providing an LOQ of approximately 126 $\mu\text{g}/\text{kg}$ shellfish (see Table 2011.27G).

(b) *Addition of samples and standards.*—Add in the following order to each of the 96 wells: 35 μL assay buffer; 35 μL STX standard, QC check, or sample extract; 35 μL [^3H] STX; 105 μL membrane preparation. The assay buffer is added first in order to wet the filter membrane. It is critical to continuously mix the membrane preparation by careful up-and-down pipetting immediately prior to dispensing into the 96-well plate to maintain an even suspension across the entire plate. Cover and incubate plate at 4°C for 1 h.

(c) *Assay filtration.*—Attach the vacuum manifold to the vacuum pump with an in-line side arm flask to catch filtrate from the plate filtration process. Set the vacuum pressure gauge on the pump or vacuum manifold to 4–8” Hg (135–270 millibar), as specified in the instructions provided with the filtration plates. Place the 96-well plate on the vacuum manifold. Fill empty wells with 200 μL MOPS/choline chloride buffer to

Table 2011.27C. Performance of individual laboratories on blind duplicates (values are in µg STX diHCl equiv./kg)

Lab	ID	Day 1	Day 2	Mean	s _r	RSD _r , %
1	MLV05	370	580	475	148	31.3
	MLV06	1100	1290	1195	134	11.2
	MLV07	1260	1010	1135	177	15.6
	MLV09	860	810	835	35	4.2
	MLV11	270	430	350	113	32.3
Avg.						18.9
2	MLV05	605	670	638	46	7.2
	MLV06	1340	1520	1430	127	8.9
	MLV07	1540	1530	1535	7	0.5
	MLV09	680	1190	935	361	38.6
	MLV11	370	350	360	14	3.9
Avg.						11.8
3	MLV05	620	250	435	262	60.1
	MLV06	1320	1460	1390	99	7.1
	MLV07	1220	1303	1262	59	4.7
	MLV09	950	1130	1040	127	12.2
	MLV11	480	460	470	14	3.0
Avg.						17.4
4	MLV05	410	430	420	14	3.4
	MLV06	1440	970	1205	332	27.6
	MLV07	1980	1000	1490	693	46.5
	MLV09	870	810	840	42	5.1
	MLV11	340	280	310	42	13.7
Avg.						19.2
5	MLV05	690	910	800	156	19.4
	MLV06	1260	1790	1525	375	24.6
	MLV07	1760	1720	1740	28	1.6
	MLV09	980	1630	1305	460	35.2
	MLV11	640	550	595	64	10.7
Avg.						18.3
6	MLV05	1070	700	885	262	29.6
	MLV06	1720	2520	2120	566	26.7
	MLV07	1530	1860	1695	233	13.8
	MLV09	1120	1390	1255	191	15.2
	MLV11	490	620	555	92	16.6
Avg.						20.4
7	MLV05	630	880	755	177	23.4
	MLV06	2090	1240	1665	601	36.1
	MLV07	1750	1150	1450	424	29.3
	MLV09	1460	1830	1645	262	15.9
	MLV11	230 ^a	1150 ^a			
Avg.						26.2
8	MLV05	660	940	800	198	24.7
	MLV06	2130	870	1500	891	59.4
	MLV07	1210	2150	1680	665	39.6
	MLV09	820	1120	970	212	21.9
	MLV11	600	410	505	134	26.6
Avg.						34.4

Table 2011.27C. (continued)

Lab	ID	Day 1	Day 2	Mean	s _r	RSD _r , %
9	MLV05	330	300	315	21	6.7
	MLV06	890	1250	1070	255	23.8
	MLV07	840	890	865	35	4.1
	MLV09	590	870	730	198	27.1
	MLV11	110	250	180	99	55.0
Avg.						23.3
Overall avg.						22.2

^a Outlier; not used in calculations.

ensure even vacuum pressure and filtration across the plate. Turn on vacuum. Optimum vacuum will pull the wells to dryness in 2–5 s. Pull contents of all wells through until all liquid is removed. (*Note:* Too low a vacuum will result in slow well clearance, but too high will result in an airlock and no well clearance.) With vacuum pump running, quickly rinse each well twice with 200 µL ice cold MOPS/choline chloride buffer using multichannel pipet. Maintain vacuum until liquid is removed.

(d) *Preparation of the assay for counting.*—Remove the plastic bottom from the plate. Blot the bottom once on absorbent toweling.

(1) *For counting in microplate scintillation counter.*—Place the microplate in a counting cassette. Seal the bottom of the 96-well plate with sealing tape. Add 50 µL Optiphase scintillation cocktail per well using multichannel pipet. Seal the top of the plate with sealing tape. Allow to incubate 30 min at room temperature. Place the plate in a counting cassette and count in a microplate scintillation counter for 1 min/well.

(2) *For counting in traditional scintillation counter.*—Place the microplate in the MultiScreen punch system apparatus. Place the disposable punch tips on top of the microplate. Punch the filters from the wells into scintillation vials and fill with 4 mL scintillation cocktail (Scintiverse or equivalent). Place caps on the vials and vortex. Allow vials to sit overnight in the dark, then count using a tritium window in a traditional scintillation counter.

G. Analysis of Data

For assays performed using the traditional counter, curve fitting is performed using a four-parameter logistic fit, also known as a sigmoidal dose response curve (variable slope; see Figure 2011.27), or Hill equation:

$$y = \min + \frac{\max - \min}{1 + 10^{(x - \log_{10} EC_{50}) \text{ Hill slope}}}$$

where max is the top plateau representing maximum binding in CPM in the absence of competing nonradiolabeled STX, also known as B₀; min is the bottom plateau, equal to nonspecific binding (in CPM) in the presence of saturating nonradiolabeled toxin; IC₅₀ is the inhibitory concentration at which CPM are 50% of max-min (dashed lines; Figure 2011.27); Hill slope is the slope of the curve; x axis is the log concentration of STX; and y axis is total ligand binding in CPM (here represented as B/B₀, or bound/max bound). A curve fitting package such as Prism (GraphPad Software, Inc.) is recommended. For the microplate counter users, receptor

Table 2011.27D. Calibration curve and QC check parameters in three receptor binding assays performed in nine participant laboratories

Lab	Assay day	Slope	IC ₅₀ , nM	QC, nM	Reference, CPM	IC ₇₀ , nM	Standards where RSD >30%; action	Curve fitting software	Scintillation counter	Manual/microplate
1	1	-0.9	1.9	2.4	720	0.90	None	Prism v 3.02	Packard Top Count	Microplate
	2	-1.0	2.0	2.6	733	0.96	None			
	3	-1.1	2.1	3.2	1038	0.92	None			
2	1	-1.1	1.8	3.8	1160	0.66	3 nM; 1 well removed	Prism v 5.0	Packard Top Count	Microplate
	2	-1.2	2.2	3.9	1260	0.85	None			
	3	-1.0	1.6	3.2	1262	0.46	3 nM, 1 nM removed			
3	1	-1.0	2.0	2.3	2529	0.41	First column removed	Prism v 5.0	Wallac Microbeta	Microplate
	2	-0.9	2.0	2.5	1463	0.92	1000 nM; 1 well removed			
	3	1.0	1.6	2.8	2088	0.80	None			
4	1	-0.9	1.7	3.4	1125	0.61	None	Prism v 3.03	PerkinElmer Tricarb	Manual
	2	-1.2	1.7	3.2 ^a	1611	0.77	None			
	3	-0.9	1.2	2.9	1324	0.45	30 nM 35%; 1 well removed			
5	1	-0.9	1.4	3.3	1566	0.64	1.0 nM; 1 well removed	MultiCalc	Wallac Microbeta	Microplate
	2	-1.2	1.8	3.6	1528	1.05	0.1 nM and 30 nM; 1 well removed			
	3	-1.2	1.8	2.9	1052	0.67	None			
6	1	-1.1	2.6	3.0	670	1.15	None	Prism v 4.0	Wallac Microbeta	Microplate
	2	-1.0	2.0	4.0 ^b	1124	1.08	None			
	3	-1.1	3.4	6.5 ^b	1030	2.04 ^c	None			
7	1	-0.8	1.0	2.8 ^a	919	0.33	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	1.6	2.7	619	0.70	None			
	3	-0.9	2.1	3.2 ^a	693	0.82	None			
8	1	-1.2	1.7	3.7	1146	0.86	None	Prism	Wallac Microbeta	Microplate
	2	-1.1	1.4	1.5 ^b	1095	0.78	None			
	3	-1.1	2.4	2.3	886	1.04	None			
9	1	-1.0	2.2	4.0 ^b	1363	0.97	None	Prism	Wallac Microbeta	Microplate
	2	-1.0	2.0	3.2	1380	0.85	100 nM 33%; left in			
	3	-1.0	2.1	3.7	1532	0.92	None			

^a One well removed.

^b Outside of specifications.

^c Outlier by Grubbs test.

assay applications provided by the manufacturer may be used (e.g., MultiCalc; PerkinElmer Wallac, Gaithersburg, MD).

(a) *Sample quantification.*—Sample quantification is carried out only on dilutions that fall within B/B₀ of 0.2–0.7, where B represents the bound [³H]STX (in CPM) in the sample and B₀ represents the max bound [³H]STX (in CPM). Where more than one dilution falls within B/B₀ of 0.2–0.7 on the curve, all sample wells corresponding to these dilutions are used to calculate sample concentration. Sample concentration is calculated in µg STX diHCl equiv./kg shellfish, from the in-well nM concentration obtained from the curve fitting software using the following formulas:

$$(nM \text{ STX equiv}) \times (\text{sample dilution}) \times \frac{(210 \mu\text{L total volume})}{35 \mu\text{L sample}} = nM \text{ STX equiv in extract}$$

$$(nM \text{ STX diHCl equiv. in extract}) \times \frac{1 \text{ L}}{1000 \text{ mL}} \times \frac{372 \text{ ng}}{\text{nmol}} \times \frac{1 \mu\text{g}}{1000 \text{ ng}} = \mu\text{g STX diHCl equiv./mL}$$

$$\mu\text{g STX diHCl equiv./mL} \times \frac{\text{mL extract}}{\text{g shellfish}} \times \frac{1000 \text{ g}}{\text{kg}} = \mu\text{g STX diHCl equiv./kg}$$

H. Assay Performance Standards

The following criteria must be met for assay acceptance:

Table 2011.27E. Results of the receptor binding assay (RBA), mouse bioassay (MBA), and HPLC analyses of 21 shellfish extracts, sorted by mouse bioassay value (all values are in µg STX diHCl equiv./kg shellfish tissue; results in bold indicate toxicity above the 800 µg STX diHCl equiv./kg regulatory limit; all other results indicate toxicity below the regulatory limit)

Sample	Lab 1	Lab 2	Lab 3	Lab 4	Lab 5	Lab 6	Lab 7	Lab 8	Lab 9	RBA, avg.	HPLC	MBA
21	ND ^a	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
5	180	200	200	150	150	100	150	290	100	168	108	ND
15	330	270	410	180	590	680	370	1570^b	90	365	196	182
13	270	370	480	340	640	290	240	600	110	371	236	299
20	430	350	460	280	550	490	1150^b	410	250	403	236	299
14	400	1240^b	560	450	650	530	500	440	200	466	625	343
1	370	610	620	410	690	1070^b	630^b	660	330	599	413	387
16	580	670	250	430	910	700	860^b	940^b	300	627	413	387
3	80	190	140	90	130	160	230	220	100	149	341	405
6	950	940	1060	1130	1040	750	1460	1320	810	1051	618	485
7	660	930	1080	870	840	1320	1490	2420	490	960	685	528
2	1100	1340	1320	1440	1260	1720	2080	2130	890	1476	931	595
17	1290	1520	1460	970	1800	2520	1470	870	1250	1460	931	595
4	860	680	950	870	980	1120	1460	820	590	926	1070	653
12	810	1190	1130	810	1630	1390	1880	1120	870	1203	1070	653
11	1260	1540	1220	1980	1760	1530	1660	1210	840	1444	965	714
18	1010	1600	1390	1000	1720	1860	1520	2150	890	1452	965	714
8	1360	1520	1580	1110	1700	3180	1400	2780	520	1683	894	752
9	830	1180	1130	1150	1130	1780	1340	980	690	1134	802	792
19	1640	2130	2800	2660	2330	1850	3390	2740	1830	2374	2000	1027
10	2440	2840	2910	1740	2150	1800	2690	2490	1210	2252	1890	1080

^a ND = Not detected.

^b Outlier; not used in average calculation.

(a) For a ligand that specifically binds at one receptor site, the slope of the resulting competition curve should theoretically be -1.0. If the slope of the curve for a given assay is outside of the acceptable range of -0.8 to -1.2, linearity of the assay will be compromised and quantification of the unknowns will be incorrect.

(b) RSDs of triplicate CPMs for standards should be below 30% as variability may affect the slope calculation and thereby quantification of samples.

(c) If the IC₅₀ is out of the acceptable range (2.0 nM ± 30%) then the assay should be considered suspect and rerun, as a shift in the curve will result in over- or underestimation of sample concentrations.

(d) QC check should be 3 nM STX ± 30% (in-well concentration). Assays with a QC check sample out of specifications should trigger a check of the IC₅₀ value.

The following criteria must be met for acceptability of a sample measurement:

(a) Sample quantification should be done only on dilutions that fall within B/B₀ of 0.2–0.7. In the event that all sample dilutions fall below B/B₀ 0.2 (i.e., concentration is too high), further dilutions must be made and the sample reanalyzed. In the event that the sample concentration is too low to be quantified (i.e., B/B₀ > 0.7), the sample is reported as below LOD. If more

Table 2011.27F. Dilution series to prepare bulk solutions for standard curve

	Stock, M	In-assay, M
100 µL 268.8 µM STX + 4.38 mL 0.003 M HCl	6 × 10 ⁻⁶	1 × 10 ⁻⁶
500 µL 6 × 10 ⁻⁶ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁷	1 × 10 ⁻⁷
1.5 mL 6 × 10 ⁻⁷ M + 3.5 mL 0.003 M HCl	1.8 × 10 ⁻⁷	3 × 10 ⁻⁸
500 µL 6 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁸	1 × 10 ⁻⁸
500 µL 1.8 × 10 ⁻⁷ M + 4.5 mL 0.003 M HCl	1.8 × 10 ⁻⁸	3 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁸ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻⁹	1 × 10 ⁻⁹
500 µL 6 × 10 ⁻⁹ M + 4.5 mL 0.003 M HCl	6 × 10 ⁻¹⁰	1 × 10 ⁻¹⁰
5 mL 0.003 M HCl	0	Reference

Table 2011.27G. Recommended microplate layout for ease of handling triplicate wells of standard curve, QC check sample, and unknown samples; each sample is run at three dilutions (1:10, 1:50, 1:200); standard curve is run in columns 1–3 (values are in M STX)^a

Microplate row	Microplate column											
	1	2	3	4	5	6	7	8	9	10	11	12
A	10 ⁻⁶	10 ⁻⁶	10 ⁻⁶	QC	QC	QC	U3	U3	U3	U6	U6	U6
							1:50	1:50	1:50	1:10	1:10	1:10
B	10 ⁻⁷	10 ⁻⁷	10 ⁻⁷	U1	U1	U1	U3	U3	U3	U6	U6	U6
				1:10	1:10	1:10	1:200	1:200	1:200	1:50	1:50	1:50
C	3 × 10 ⁻⁸	3 × 10 ⁻⁸	3 × 10 ⁻⁸	U1	U1	U1	U4	U4	U4	U6	U6	U6
				1:50	1:50	1:50	1:10	1:10	1:10	1:200	1:200	1:200
D	10 ⁻⁸	10 ⁻⁸	10 ⁻⁸	U1	U1	U1	U4	U4	U4	U7	U7	U7
				1:200	1:200	1:200	1:50	1:50	1:50	1:10	1:10	1:10
E	3 × 10 ⁻⁹	3 × 10 ⁻⁹	3 × 10 ⁻⁹	U2	U2	U2	U4	U4	U	U7	U7	U7
				1:10	1:10	1:10	1:200	1:200	1:200	1:50	1:50	1:50
F	10 ⁻⁹	10 ⁻⁹	10 ⁻⁹	U2	U2	U2	U5	U5	U5	U7	U7	U7
				1:50	1:50	1:50	1:10	1:10	1:10	1:200	1:200	1:200
G	10 ⁻¹⁰	10 ⁻¹⁰	10 ⁻¹⁰	U2	U2	U2	U5	U5	U5			
				1:200	1:200	1:200	1:50	1:50	1:50			
H	REF	REF	REF	U3	U3	U3	U5	U5	U5			
				1:10	1:10	1:10	1:200	1:200	1:200			

^a REF = Reference; QC = quality control check; U = unknown sample. [Note: The same standard curve may be used for multiple plates (i.e., 11 samples can be run on subsequent plates in a series if the standard curve is not included).]

than one dilution falls on the linear part of the curve, an average value calculated from all dilutions should be used. If there is disagreement between different dilutions in final concentration reported, check for error in the sample dilution process.

(b) RSD of the sample CPMs should be ≤30%.

Reference: *J. AOAC Int.* **95**, 795(2012)

Results and Discussion

Sample Characterization

All shellfish homogenates (MLV1–16) were analyzed by

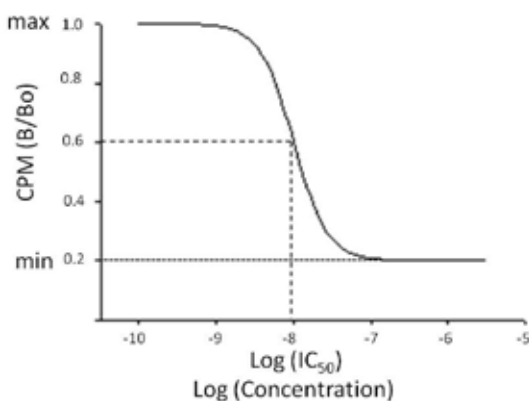


Figure 2011.27. Sigmoidal dose response curve. Dashed lines indicate log IC₅₀.

HPLC using the precolumn oxidation method (AOAC Method **2005.06**) to determine toxin congener profiles and quantify total PST as µg STX diHCl equiv./kg prior to initiation of the study (Table 2). It is noteworthy that the clear majority of samples, irrespective of shellfish species and location, were dominated largely by STX and GTX2,3 whereas the N1-hydroxylated congeners NEO and GTX1,4 were virtually absent, except in blue mussel from the U.S. west coast. The most unusual profile was observed in green mussel, which was dominated by the weakly toxic N-sulfo-carbamoyl congeners C1,2. The samples were analyzed by the AOAC mouse bioassay (AOAC Method **959.08**) by three laboratories that routinely perform the mouse bioassay for regulatory purposes (Table 3). The mouse bioassay detection limit is approximately 400 µg STX diHCl equiv./kg (one laboratory reported values as low as 290 µg STX equiv./kg). Because the study design included samples that bracketed the lower regulatory limit of 400 µg STX diHCl equiv./kg, several samples were reported as being below the mouse bioassay detection limit. For samples in which all values were above the detection threshold, the between-laboratory RSD_R of the mouse bioassay was 18.9%.

Data Reporting and Initial RBA Data Review

Nine of the 10 laboratories that received the study materials completed the study and reported results. All nine carried out the practice assay and reported results to the coordinating laboratory, which evaluated the results and provided feedback to the participating laboratories before initiating the full study. Following completion of the full study, the participating laboratories provided all raw and calculated data for each of

the three assays performed via e-mail to the coordinating laboratory. The calculated results sheets were reviewed by the coordinating laboratory for obvious errors in sample dilutions and calculations, and for the use of the prescribed curve-fitting model. One laboratory used a sigmoidal curve-fitting model with the slope set to 1 (one-site binding curve in Prism), rather than the prescribed four-parameter logistic fit. In this case, the raw data were reanalyzed by the coordinating laboratory using the prescribed method. Obvious errors in calculation were corrected, such as accounting for the two-fold sample dilution resulting from the extraction process. In some cases, the participating laboratory carried out a fourth assay due to variability or inconsistency among dilutions for selected samples. In these cases, the value reported from the repeat (fourth) assay was used. One laboratory had consistent disagreement between the 1/50 and 1/200 dilutions when both fell within B/B_0 , 0.2–0.7. In all cases the 1/200 dilution overestimated almost two-fold relative to the 1/50 dilution, suggesting a systematic dilution error. In standard practice, these samples should be rerun. However, the instructions did not direct the participants to do so. Therefore, where there was corroborative evidence for the value reported by the 1/50 dilution, based on the 1/10 dilution, the 1/200 dilution was omitted. Where there was no basis on which to exclude the 1/200 value, an average value was calculated. This tended to result in an overestimate, and in two cases resulted in statistical outliers.

Overall Performance of the Method: Reproducibility

Table 2011.27A summarizes the results obtained for 21 individual shellfish samples analyzed in three RBAs, determined by nine participating laboratories. Samples 1–7 were analyzed in the first assay, samples 8–14 in the second assay, and samples 15–21 in the third assay. Among these samples were five blind duplicates, treated here as individual unknown samples. One sample (marked by an footnote a in Table 2011.27A) had a high variability in CPM between wells that was not attributable to any known cause, and was, therefore, omitted from analysis. Outliers identified by Grubbs test ($P < 0.01$) were excluded from the analysis (marked by footnote b in Table 2011.27A). The overall RSD_R among all 21 independent samples was 33.2%, resulting in an average HorRat value of 2.0 (Table 2011.27A). The HorRat values on individual samples ranged from 1.4 to 3.3, with a median value of 1.8. There was no apparent trend in reproducibility according to sample concentration or among shellfish species. If only the laboratories that are routine users of the RBA for PST (Laboratories 1–5) are included in the analysis, the average RSD_R is 23.1%, resulting in an average HorRat value of 1.4. Laboratory 9 tended to report the lowest values among the participating laboratories (14 of 21 samples), and although its individual sample values were not found to be statistical outliers, removing the results of this laboratory reduces all but one HorRat value (which remains unchanged), yielding an average HorRat value of 1.8 (range 1.0–2.8; Table 2011.27A). Removal of any other single laboratory's results does not appreciably change the overall study performance. The reason for the systematically low values reported by Laboratory 9 is not clear, since the assay parameters fall well within those reported by the other laboratories. Given that assay parameters are within normal range, one possible source of systematic error

could be incomplete extraction or pH adjustment of extracts, either of which would result in lower toxicity values.

A comparison of the RBA reproducibility with that of existing AOAC *Official Methods* is instructive. The AOAC collaborative study of the mouse bioassay (11), which entailed the analysis of seven samples representing three levels of STX-spiked shellfish by 11 participating laboratories, yielded a similar average RSD_R of 22%. More recent proficiency tests of the mouse bioassay performed in European regulatory laboratories report RSD_R of 2.3–38.3% on three samples run by eight laboratories (2) and RSD_R of 18.1–44.8% on two samples run by 20 laboratories (12). The mouse bioassay RSD_R values obtained in the current study ranged from 1.1 to 46.3% (average 19%) for three laboratories. The collaborative studies of the HPLC methods report reproducibility values for individual PST congeners, but do not report reproducibility of the composite toxic potency values. Collaborative study of the precolumn oxidation HPLC method (AOAC Method 2005.06) resulted in an average RSD_R of 27.0% and HorRat value of 1.3 (range 0.8–2.1) for STX following C18 cleanup, but the reproducibility of other congeners varied considerably, with the maximum HorRat value (4.7), exceeding the highest HorRat value obtained by RBA (3.3).

Because composite toxic potency values were not reported in the studies of the HPLC methods, it is uncertain how this variability influences the composite toxic potency calculated from these methods. The average and ranges of HorRat values obtained for different congeners were: neoSTX–1.7 (range 1.2–2.5); dcSTX–1.1 (range 0.6–2.1); GTX1,4–1.9 (range 1.1–4.2), GTX2,3–1.4 (range 0.8–1.9); B1–1.1 (range 0.7–1.9); and C1,2–1.6 (range 0.9–4.5). Because of the variability obtained in neoSTX, GTX1,4, C3,4, and B2, AOAC Method 2005.06 calls for a second SPE-COOH cleanup of samples suspected of containing these congeners, after which reproducibility improved somewhat: neoSTX–1.8 (range 1.3–2.1); GTX1,4–1.3 (range 1.0–2.1); and C3,4–1.2 (range 0.8–1.8). The postcolumn oxidation HPLC method (AOAC Method 2011.02) reported an average HorRat value of 0.6 for STX. In this method, neoSTX with an average HorRat of 1.9 (range 0.6–4.0) and GTX4 with an average HorRat of 1.6 (range 1.0–2.9) had reproducibility values that may affect the overall composite potency values. The maximum HorRat value (4.0) reported in this study also exceeded the maximum value reported in the RBA.

In summary, with the removal of Laboratory 9, the overall reproducibility of the RBA falls within the performance measures achieved by the established AOAC *Official Methods* for PST. The difference in reproducibility achieved by the laboratories that are routine users of the assay and participants who are not routine users of the method highlights the importance of training if this method were to be implemented in a regulatory setting.

Within-Laboratory Repeatability

Within-laboratory variability (RSD_r) was determined on five samples that were provided as blind duplicates. Participants were unaware that blind duplicates were included among the coded samples received. The duplicate samples were coded so that they were analyzed in separate assays, with different duplicate pairs falling into different assays (Table 1). One outlier was found among the results of the blind duplicates by Cochran's

test, $P < 0.025$ (Laboratory 7, sample MLV11) using the AOAC INTERNATIONAL Interlaboratory Study Workbook for Blind Duplicates, v2.0. An overall RSD_r of 25.1% was observed, with an RSD_R of 32.9%, yielding a HorRat value of 2.0, similar to that of the overall study (Table 2011.27B). When the performance of individual laboratories was evaluated separately, the average RSD_r was 22.2%, with individual laboratories varying from 11.8 to 34.4% (Table 2011.27C). Routine users of the microplate format of the PST RBA (Laboratories 1–5) obtained an average RSD_r of 17.1%, which is similar to that obtained in the SLV study (10), and lower than that obtained by nonroutine users (Laboratories 6–9), which averaged 26.1% and ranged as high as 34.4%. The AOAC collaborative study of the mouse bioassay (11) did not report RSD_r ; however, analysis of the data from that study using AOAC INTERNATIONAL's Interlaboratory Study Workbook for Blind Duplicates results in an average RSD_r of 16.5% for three STX-spiked samples. Proficiency testing of the mouse bioassay performed in eight French laboratories reported an average RSD_r of 8.3% on three samples (2). The analysis of blind duplicates in the collaborative study of the precolumn oxidation HPLC method (AOAC Method 2005.06) resulted in an RSD_r of 15.2% for STX following SPE C18 cleanup and an average RSD_r of 16.4% across all congeners, which ranged from 6.0 to 31.7%. Following SPE-COOH cleanup, repeatability was similar, with RSD_r of 17.2% across all congeners. The intralaboratory repeatability values obtained in the postcolumn oxidation HPLC method (AOAC Method 2011.02) averaged 6.4% for STX; most other congeners were similar, with neoSTX being the only congener that showed a somewhat higher RSD_r of 23.3%.

In summary, the within-laboratory repeatability of the RBA was found to be acceptable, with all but two laboratories achieving an RSD_r of 23.3% or less, and the routine users of the assay achieving an average RSD_r of 17.1%.

Spike Recovery

Three samples included in the study were homogenates of blue mussel spiked with STX diHCl at concentrations intended to bracket the regulatory limits of 800 μg STX equiv./kg used by most countries and 400 μg STX equiv./kg imposed in the Philippines. Nominal concentrations in the spiked samples were 200, 500, and 1200 μg STX equiv./kg. Also included in the study was the blue mussel homogenate to which the STX spikes had been added, which was determined to be negative for STX by the precolumn oxidation HPLC method. The negative control homogenate was reported as nondetectable by eight of nine laboratories. Recovery of spiked STX by the RBA was 84.4, 93.3, and 88.1%, respectively, for the 200, 500, and 1200 μg STX diHCl equiv./kg spike levels, and yielded a slope of 0.87 and r^2 of 0.86 (Figure 2). In the current study, the mouse bioassay reported < detection limit, and 68.6 and 40.5% recovery for the 200, 500, and 1200 μg STX diHCl equiv./kg spike levels. The AOAC collaborative study of the mouse bioassay (11) reported recoveries of 62.3% at spike levels similar to those in the current study (equivalent to 1000 μg STX diHCl equiv./kg) but higher recoveries of 81.5 and 96.0% were achieved at higher spike levels equivalent to 4000 and 8000 μg STX diHCl equiv./kg.

The observed poor recovery in the mouse bioassay at concentrations near and below the regulatory limit has been observed in other studies (2), and has been attributed to a

salt or protective effect of the shellfish matrix, which, for concentrations at or below the regulatory limit of 800 $\mu\text{g}/\text{kg}$, is injected undiluted into the mouse. The spike recovery observed in the precolumn HPLC method in this study is also somewhat low, with 54.0, 62, and 51.5% recovery at the 200, 500, and 1200 μg STX diHCl equiv./kg spike levels, respectively. The AOAC collaborative study of the precolumn HPLC method reported 74.4–76.8% at similar spike levels following SPE C18 cleanup and 63.7–68.2% following SPE-COOH cleanup (3, 4). In comparison, the postcolumn HPLC method reported 88–104% recovery of STX spiked at levels somewhat lower than the current study. The higher recovery of the RBA than the HPLC method in the current study may reflect the use of the 0.1 M HCl extraction method in the RBA as compared to the acetic acid extraction used in the HPLC methods.

We previously established in the SLV study that the RBA performs well with shellfish extracted using either method (10). In that study, the RBA reported slightly higher toxicity values for shellfish extracts made using the 0.1 M HCl method than the acetic acid extraction, yielding a correlation of 0.99 with a slope of 1.23 (10). The higher toxicity reported by the RBA in 0.1 M HCl extracts may reflect the hydrolysis of less toxic congeners to more toxic congeners.

Assay Parameters and Quality Metrics

Table 2011.27D summarizes the assay parameters and quality metrics for all laboratories. Eight of nine laboratories used microplate scintillation counters. Laboratory 4 used the manual counting method in which the microplate well filters are punched out, using an eight-place punch system, into traditional 4 mL scintillation vials and counted. Its performance using the manual counting method (RSD_r 17.4%) was similar to or better than that of the laboratories using the microplate method, indicating that using the manual counting method does not affect the performance of the assay. Similarly, there was no apparent difference in assay parameters when the Packard Top Count (single detector) was used, compared to the Wallac Microbeta (coincidence detector), although the reference CPM values obtained on the Top Count generally were somewhat lower due to differences in counting efficiency inherent in the differences in detector geometry. Eight of nine laboratories used GraphPad Prism for curve-fitting, while only Laboratory 5 used Wallac MultiCalc software. Values reported by Laboratory 5 fell well within the range of values reported by laboratories using Prism.

All assays resulted in slopes between -0.8 and -1.2 , as specified in the protocol. This specification reflects the fact that in a competitive binding assay for a ligand that interacts specifically at a single receptor site, the slope of the resulting standard curve should theoretically be 1.0. Although curve-fitting software packages often include a one-site binding curve that fixes the slope at 1.0, we specified in the protocol the use of the four-parameter logistic fit (also known as sigmoidal dose-response with variable slope), because it more readily identifies problems with the standard curve that may skew results. Laboratory 9 reported results using a one-site binding curve fit; in this case, the coordinating laboratory recalculated their raw data using the four-parameter logistic fit. The protocol also calls for $RSD\% < 30$ on all standards. Most analysts did not experience variability problems in the standard wells. Infrequent high RSD s were most often associated with the well

Table 2. Congener profiles in shellfish homogenates included in the collaborative study^a

Sample name	Species	STX	NEO	dcSTX	GTX1,4	GTX2,3	dcGTX2,3	B1	C1,2	C3,4	Total PSP	µg STX diHCl equiv./kg
MLV01	Surf clam	639.8		74.0		226.2	207.0				1146.9	894.3
MLV02	Almeja clam	298.3				1290.1		266.6			1855.0	802.1
MLV03	Chorito mussel	77.6				310.4					388.0	195.5
MLV04	Atlantic sea scallop	831.6				2785.6					3617.3	1890.2
MLV05	Atlantic sea scallop	193.8				576.2					770.0	412.8
MLV06	California mussel	912.8		10.9		0.0		233.8			1157.5	931.3
MLV07	Blue mussel, U.S. east coast	548.2				1097.3					1645.5	965.2
MLV08	Green mussel	164.2		63.5			272.3	454.8	3629.0		4419.6	340.8
MLV09	Blue mussel, U.S. west coast	432.3	124.9	8.7	353.7	727.8		506.4			2153.9	1070.9
MLV10	Butter clam	1763.5		40.6		533.2		203.5			2540.8	2000.9
MLV11	Almeja clam	159.1		12.2		185.5					356.8	236.9
MLV12	Blue mussel spike	108.4									108.4	108.4
MLV13	Blue mussel spike	310.2									310.2	310.2
MLV14	Blue mussel spike	618.5									618.5	618.5
MLV15	Blue mussel blank										0.0	0.0
MLV16	Chorito mussel	389.8		14.3		754.1					1158.1	684.9

^a Values for individual congeners are in µg/kg. Values for composite toxicity are in µg STX diHCl equiv./kg. Abbreviations for congeners are as follows: STX – saxitoxin; NEO – neosaxitoxin; dcSTX – decarbamoyl saxitoxin; GTX1,4 – gonyautoxin 1 and gonyautoxin 4; GTX2,3 – gonyautoxin 2 and gonyautoxin 3; B1 – gonyautoxin 5 (also known as sulfocarbamoyl STX B1); C1,2 – sulfocarbamoyl STX C1 and sulfocarbamoyl STX C2; C3,4 – sulfocarbamoyl STX C3 and sulfocarbamoyl STX C4.

Table 3. Mouse bioassay results on collaborative study samples from three laboratories^a

Sample No.	Sample ID	MBA Lab A	MBA Lab B	MBA Lab C	MBA Avg.	MBA s _R	MBA RSD _R , %
1	MLV05	400	415	340	385	39.7	10.3
2	MLV06	550	597	540	562	30.4	5.4
3	MLV08	440	<dl ^b	370	405	49.5	12.2
4	MLV09	670	612	760	681	74.6	11.0
5	MLV12	<dl	<dl	<dl	—	—	—
6	MLV14	489	489	480	486	5.2	1.1
7	MLV16	585	585	470	547	66.4	12.1
8	MLV01	750	716	600	689	78.6	11.4
9	MLV02	670	1115	590	792	282.9	35.7
10	MLV04	2040	<dl	1080	1560	678.8	43.5
11	MLV07	1480	748	670	966	446.8	46.3
12	MLV09	—	594	670	602	11.3	1.9
13	MLV11	380	379	<dl	380	—	—
14	MLV13	<dl	343	<dl	343	—	—
15	MLV03	400	364	<dl	382	—	—
16	MLV05	—	396	370	383	18.4	4.8
17	MLV06	—	702	630	666	50.9	7.6
18	MLV07	—	<dl	690	690	—	—
19	MLV10	1320	890	870	1027	254.2	24.8
20	MLV11	—	364	290	327	52.3	16.0
21	MLV15	<dl	<dl	<dl	—	—	—

^a Values are in µg STX diHCl equiv./kg.

^b dl = Detection limit.

in column 1 of the 96-well plate. Most analysts removed the suspect well from the curve-fitting process. When the RSD for a given standard was near the stated cutoff (e.g., 31–33%), and left in the curve-fitting process, there was no apparent effect on the curve parameters listed as criteria for assay acceptance.

The average IC₅₀ among all 27 assays was 1.9 + 0.45 nM (RSD_R 23.5%). The other assay quality metric called for by the protocol is the analysis of the QC check sample, which should be 3 ± 0.9 nM STX (30% RSD, in-well concentration). Four of the 27 assays had QC values outside the stated limits, with no obvious error responsible for the variability. Among these, Laboratory 7 reported 6.5 nM for the QC check in assay 3 and an IC₅₀ of 3.4 nM, which was outside the norm. Similarly, Laboratory 8 reported a QC of 1.5 nM in assay 2 and a low IC₅₀ of 1.4 nM, which is at the lower edge of acceptability. In general practice, these values would trigger repeating the assay. However, because of the minimal number of laboratories participating in the study, both of these assays were retained in the study. In neither case were the reported sample values systematically higher or lower than those reported in the other assays.

LOD and LOQ

The LOD was calculated based on the measurement of the negative control shellfish matrix (MLV15) using the blank + 3×SD approach according to Eurachem guidelines (13), as

recently applied to AOAC Method **2006.02**, an ELISA for domoic acid in shellfish using a similar four-parameter logistic curve (14). All laboratories reported <dl for this sample using the prescribed cutoff of B/B₀ <0.7 for quantification, with the exception of Laboratory 8, which was removed as an outlier as determined by Grubbs test (*P* < 0.01). If these samples are instead quantified using the B/B₀ values obtained, a mean of 5.5 ng/mL is obtained with an SD of 5.7 ng/mL, resulting in an LOD of 45 µg STX diHCl equiv./kg. Using the blank + 10×SD definition, an LOQ of 126 µg STX di HCl equiv./kg is thus obtained. We previously established empirically that a 1/10 dilution of shellfish extracts is sufficient to remove matrix effects in the RBA (10), when a quantification cutoff of B/B₀ <0.7 is used. This is the basis for the ten-fold minimum sample dilution used in the current study. The IC₇₀ values (B/B₀ 0.7) for all standard curves run in the study are presented in Table **2011.27D**. An average of 0.80 ± 0.188 nM STX diHCl was obtained across all assays, following the removal of one outlier based on the Grubbs test (*P* < 0.01). Applying the blank + 3×SD to this value, an LOD of 64 µg STX diHCl equiv./kg is obtained; applying the blank + 10×SD to this value results in an LOQ of 131 µg STX diHCl equiv./kg for a sample diluted 1/10 and extracted as indicated in the study, in fair agreement with the value calculated above.

Correlation with HPLC and Mouse Bioassay

Comparison of the RBA results with the mouse bioassay

Nominal	Avg	S _R	RSD _R , %	Recovery, %
200	169	58	34.6	84.4
500	466	133	28.5	93.3
1200	1057	228	21.7	88.1

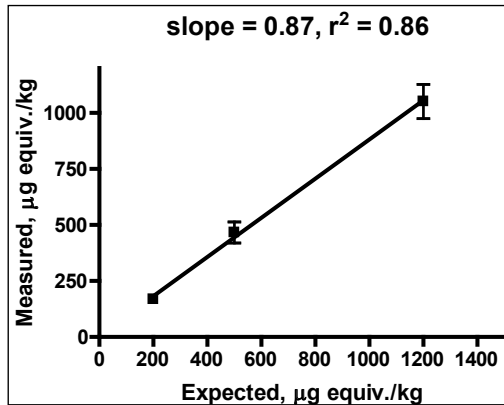


Figure 2. Recovery of spiked STX diHCl in homogenates of blue mussel. Values are in µg STX diHCl equiv./kg.

results yielded an r^2 of 0.84 and a slope of 1.64, indicating that the RBA reports somewhat higher STX equivalents in shellfish, relative to the mouse bioassay (Figure 3). This overestimate has been previously reported for both RBA and HPLC methods (2, 9) at the STX levels near or below the regulatory limit, which are the focus of the current study. Consistent with these findings, the HPLC method also reported higher values than the mouse bioassay in this study, with a slope of 1.33 and an r^2 of 0.84. RBA results correlated better with the precolumn oxidation HPLC method, with a slope of 1.20 and an r^2 of 0.92.

RBA Yielded No False Negatives Relative to the Regulatory Limit

When the data from the three methods were sorted by increasing µg STX diHCl equiv./kg as reported by the mouse bioassay, the RBA did not report any false negatives when compared to the regulatory limit of 800 µg STX equiv./kg (Table 2011.27E). When compared with the precolumn oxidation HPLC method, only Laboratory 9 reported values lower than the HPLC method. The fact that the RBA reports somewhat higher toxicity than the mouse bioassay or HPLC at levels near or below the regulatory limit is beneficial from a food safety standpoint. The higher values reported presumably arise from better recoveries, as demonstrated above. From a shellfish producer’s perspective, the improved detection limits relative to the mouse bioassay and better recovery of low toxin levels compared to the HPLC can help to provide advance warning of developing toxicity, allowing producers to harvest early, delay harvest, or move cultures, as appropriate.

Participants’ Comments

Laboratory 6 participated in the study without previous

experience running receptor assays, and in doing so, identified several points needing clarification that have since been added to the proposed *Official Method* as enumerated in this report: (1) The vacuum required for filtration was not specified at 4–8” Hg, which is critical because insufficient vacuum pressure results in too slow a clearance of the wells, whereas too much pressure results in an airlock and no filtration at all. (2) Scintillation counting time for the microplates is 1 min/well. (3) Instructions have been added regarding how to calculate sample concentration if more than one dilution falls within B/B₀ 0.2–0.7; specifically, an average value should be calculated from all sample dilutions falling within B/B₀ 0.2–0.7. When corrected for dilution, serial sample dilutions should yield similar quantification. The absence of linearity between sample dilutions indicates either error in dilution or sample matrix interference; however, at the minimum sample dilutions recommended in the proposed *Official Method*, matrix effects from shellfish homogenates have not been encountered (10). In the current study, the nonlinearity of dilutions experienced in several samples by Laboratory 8 was not observed by the other laboratories, suggesting a systematic sample dilution issue rather than a sample matrix problem. Although experienced in RBAs in general, Laboratory 8 had not previously run the microplate filtration format of the assay for PST.

Laboratory 9, which reported generally lower values than the other laboratories, although familiar with the assay, had not performed it in more than a year. The lower values reported do not appear to be associated with conduct of the assay, or scintillation conduct of the assay, or scintillation counting, since the assay metrics are well within the averages reported by the other laboratories. Insufficient boiling or pH adjustment of sample extracts are a possible explanation. These points identified by the study participants should be added to the critical steps identified in the SLV study (10) that can affect precision and accuracy of the assay results, including: (1) ensure that the water is strongly boiling during extraction; (2) carefully adjust pH of extracts; (3) ensure even distribution of the membrane preparation across the microplate by frequent vortex-mixing or pipetting before and during its addition to the plate; (4) the wells must clear within 2–5 s during filtration; (5) the wash buffer should be ice-cold to minimize the rate of toxin release from the receptor; and (6) following addition of scintillant to the wells, incubate a minimum of 30 min to ensure that the scintillant fully penetrates the filters before counting.

Recommendations

The collaborative study of the RBA for PST was completed by nine laboratories representing six countries. Collaborators quantified PST as a composite toxicity value reported in µg STX di HCl equiv./kg in a variety of shellfish species from different regions of the world, containing varied toxin congener profiles. The study included laboratories with extensive experience as well as others with little or no previous experience. The study also included both microplate and scintillation counters as end points, because either instrument type could potentially be used by test laboratories. The study demonstrates that the RBA yields adequate repeatability, reproducibility, and recovery for routine determination and monitoring of PST in shellfish. The greater precision attained by laboratories that received prior training on the RBA and routinely implement this assay suggests that

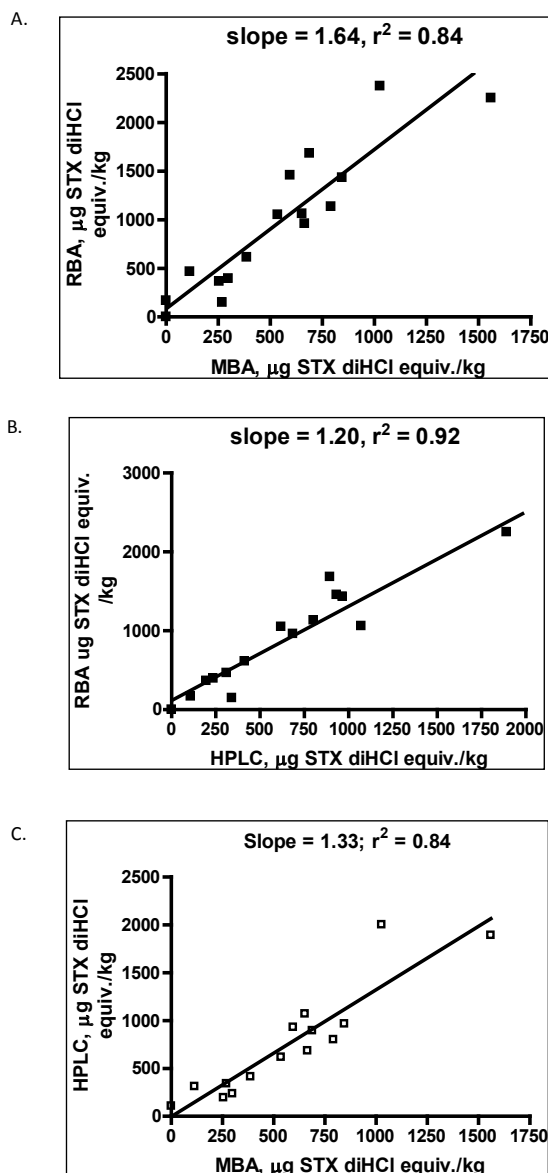


Figure 3. Correlation of the RBA results on PSP toxins in shellfish homogenates with mouse bioassay (A) and HPLC (B). Correlation between the current AOAC Official Methods, mouse bioassay, and HPLC (C).

the overall interlaboratory reproducibility can be further improved. It is recommended that this method be accepted by AOAC INTERNATIONAL as Official First Action for the determination of PST in shellfish.

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Appendix: Rat Brain Membrane Preparation

The rat brain membrane preparation used in this assay can be produced in bulk, aliquotted, and stored at -80°C until use. Under this storage condition, the preparation is stable for a minimum of 6 months. The following protocol provides sufficient membrane preparation for a minimum of 125 plates and can be scaled up or down as needed.

A. Apparatus

- (a) *Teflon/glass homogenizer*.—Motorized tapered Teflon pestle and glass tube, 15 mL.
- (b) *Motorized tissue homogenizer*.—Polytron or small hand-held blender.
- (c) *High-speed centrifuge and fixed angle rotor*.—Capable of $20\,000 \times g$ (rcf).
- (d) *Centrifuge tubes*.—12–15 mL rated for $>20\,000 \times g$ (rcf).
- (e) *Plastic cryovials*.—2 mL.
- (f) *Graduated beaker*.—300 or 500 mL.
- (g) *Pipets*.—Disposable 5 and 10 mL.
- (h) *Forceps*.

B. Reagents

- (a) *20 Rat brains*.—Male, 6-week-old Sprague-Dawley (Hilltop Lab Animals, Inc., Scottsdale, PA; <http://hilltoplabs.com>) or equivalent.
- (b) *MOPS*.—pH 7.4 (Sigma, St. Louis, MO; Cat. No. M3183-500G).
- (c) *Choline chloride*.—100 mM (Sigma; Cat. No. C7527-500G).
- (d) *Phenyl methylsulfonyl fluoride (PMSF)*.—Sigma; Cat. No. P7626.
- (e) *Isopropanol*.

C. Procedure

(1) Prepare 1 L 100 mM MOPS buffer, pH 7.4, containing 100 mM choline chloride (detailed protocol in E, above) and 0.1 mM PMSF. PMSF must first be dissolved in isopropanol; dissolve 0.174 g PMSF in 10 mL isopropanol to make 100 mM stock. Aliquot and store at -20°C . Add PMSF (1/1000, 0.1 mM final concentration) to the MOPS/choline chloride buffer fresh on the day of use.

(2) Remove medulla and cerebellum from each brain using forceps and discard. Place the cerebral cortex (see Figure 1) in a small amount of ice-cold buffer and place on ice.

(3) Place one cerebral cortex in 12.5 mL MOPS/choline Cl/PMSF, pH 7.4, in glass/teflon homogenizer (two brains in 25 mL buffer will fit into 30 mL homogenizer tube). Homogenize at 70% full speed (385 rpm) with at least 10 up and down strokes (more if necessary to homogenize brain; there should be no visible chunks remaining in the homogenate). Keep tube in ice at all times. Pour homogenized tissue into 250 mL beaker on ice and repeat procedure with remaining cortices.

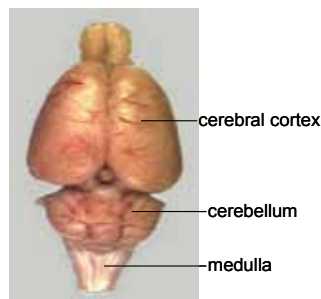


Figure 1. Rat brain.

(4) Transfer pooled homogenized tissue to centrifuge tubes, balance the tubes (pairwise; use ice-cold buffer to balance), and centrifuge at $20\,000 \times g$ for 15 min at 4°C .

(5) Aspirate the supernatant and resuspend the pellets in ice-cold MOPS/choline Cl/PMSF buffer, using an adequate amount (~5 mL) to fully resuspend the pellet (can use clean glass stir rod to break up pellet), not exceeding 10 mL per brain.

(6) Pool resuspended membrane preparation in a small beaker. Rinse centrifuge tubes with a small amount of ice-cold buffer to recover all of the membrane preparation. Bring total volume to 200 mL total (keep on ice).

(7) Keeping the beaker on ice, Polytron (or use a small hand-held blender at low speed) at 70% full speed for 20 s to obtain a consistent homogenate.

(8) Aliquot 2 mL/tube into cryovials. It is critical to keep the preparation well mixed while dispensing, e.g., prior to each aliquot to ensure equal allocation of protein/receptors to each vial. Keep cryotubes on ice.

(9) Freeze and store at -80°C . This preparation is stable for at least 6 months. Use a permanent marker to label the preparation date on the storage container.

D. Protein Assay

(a) Determine protein concentration of membrane preparation using Pierce Micro BCA Protein Assay Reagent Kit No. 23235 (microplate method) or No. 23225 (tube method) protein assay kit or equivalent protein assay (Thermo Fisher, Rockford, IL). The above protocol should yield 6–8 mg protein/mL of rat membrane preparation.

(b) Determine membrane dilution needed for the assay. The protein concentration in the daily working stock for the assay should be 1 mg/mL (this is diluted in the assay to yield 0.5 mg/mL in-assay concentration). Based on the protein concentration determined in the protein assay, determine the dilution needed to achieve 1 mg/mL. This is the dilution used in section E(e) above for all assays using this lot of membrane preparation. The protocol above typically yields a protein concentration that requires a dilution of 1/6–1/8. (Do not use less than 1/4 dilution or filtration wells may become clogged.) Protein concentration will need to be determined for each new batch of membrane preparation.

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Evaluation of variability and quality control procedures for a receptor-binding assay for paralytic shellfish poisoning toxins

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The receptor-binding assay (RBA) method for determining saxatoxin (STX) and its numerous analogues, which cause paralytic shellfish poisoning (PSP) in humans, was evaluated in a single laboratory study. Each step of the assay preparation procedure including the performance of the multi-detector TopCount[®] instrument was evaluated for its contribution to method variability. The overall inherent RBA variability was determined to be 17%. Variability within the 12 detectors was observed; however, there was no reproducible pattern in detector performance. This observed variability among detectors could be attributed to other factors, such as pipetting errors. In an attempt to reduce the number of plates rejected due to excessive variability in the method's quality control parameters, a statistical approach was evaluated using either Grubbs' test or the Student's *t*-test for rejecting outliers in the measurement of triplicate wells. This approach improved the ratio of accepted versus rejected plates, saving cost and time for rerunning the assay. However, the potential reduction in accuracy and the lack of improvement in precision suggests caution when using this approach. The current study has recommended an alternate quality control procedure for accepting or rejecting plates in place of the criteria currently used in the published assay, or the alternative of outlier testing. The recommended procedure involves the development of control charts to monitor the critical parameters identified in the published method (QC sample, EC₅₀, slope of calibration curve), with the addition of a fourth critical parameter which is the top value (100% binding) of the calibration curve.

Keywords: receptor-binding assay; paralytic shellfish poisoning; saxitoxins; variability

Introduction

Coastal regions with a history of the occurrence of paralytic shellfish poisoning (PSP) toxins present unique challenges to the organisations responsible for protecting public health. The agencies responsible for monitoring these toxins in shellfish (e.g., mussels, oysters) and other seafood species have traditionally relied on the mouse bioassay (MBA) (American Public Health Association (APHA) 1970). Until recently this live animal assay has been the only method recognised by the National Shellfish Sanitation Program (NSSP) as administered by the US Food and Drug Administration (USFDA). The MBA has served these monitoring programmes well over the decades, but the continued use of live animals for toxin testing presents practical and ethical concerns. The MBA is also recognised as having relative poor accuracy and precision due to matrix effects at low dilutions and inherent differences in response among animals. As a result there has been a considerable amount of work and progress in the development of alternative methods including a receptor-binding assay (RBA) method (Doucette et al. 1997;

Powell and Doucette 1999; Ruberu et al. 2003) and HPLC methods (Lawrence et al. 2005; van de Riet et al. 2009). The latter HPLC method has recently been accepted by the Interstate Shellfish Sanitation Conference (ISSC) and USFDA for use within the NSSP. The RBA method has recently been issued as an Official Method of Analysis (OMA) (number 2011-27) by the Association of Official Analytical Chemists (AOAC), but has yet to be presented to the ISSC for acceptance.

Most, if not all, monitoring programmes have similar requirements with respect to an acceptable replacement method for the MBA. Analytical turn-around time and sample throughput are critical factors for getting data into the hands of managers quickly, so decisions can be made regarding quarantines and notification of the public. Shellfish sample collection and shipment to an accredited laboratory can introduce significant time delays, often 24–48 h, placing the responsible agency at an immediate disadvantage in its efforts to protect consumers. Therefore, there is a need for a method that can provide data within hours of

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sample arrival. The preferred method must also be capable of automation to accommodate a significant influx of samples when PSP levels begin increasing in a region. It is also highly preferable that the methodology be simple enough to be carried out by a trained technician, as opposed to the more technically sophisticated methods that require an experienced analyst with an advanced degree. Other desirable features include improved accuracy, precision and sensitivity relative to the current MBA. The lack of precision of the MBA creates ambiguity when results are close to the action level (80 µg of saxitoxin (STX) equivalents per 100 g of shellfish tissue, abbreviated as 80 µg/100 g). Replication would help alleviate this ambiguity but is usually impractical when large numbers of samples are being assayed and throughput time must be minimised.

The van de Riet HPLC method may be the alternative method of choice for some regulatory laboratories. One seemingly minor but very important practical consideration in this regard is the administrative location of the current MBA work. If this work is presently conducted in a laboratory section in which chemical instrumental analyses are also conducted (e.g., HPLC methods), then the adoption of the HPLC method for PSP toxins could be a relatively simple transition both technically and organisationally. However, if the MBA work is currently conducted in a microbiology setting, a number of obstacles may preclude adoption of a chemical instrumental method in favour of an assay format more familiar to the microbiologist, such as an immunoassay or receptor assay. The potential obstacles in these times of reduced resources include retraining or hiring new staff, purchasing of expensive equipment or transferring resources from one department to another. Although HPLC technology includes automation via autosamplers, other factors such as time for careful filtration makes the analytical time spent per sample long enough that results for many of the samples in the queue are not available until the following workday. Furthermore, at present there are standards commercially available through the National Research Council of Canada for 12 of the more than 30 analogues of STX. The cost of these standards, and the lack of a domestic supply, may be of concern for a regulatory laboratory that processes thousands of samples per year. The detailed, compound-specific information provided by the current HPLC methods will provide valuable insight into the toxin profile(s) present along a coastal region, but may not be essential for routine monitoring purposes. A quick and reliable estimate of total toxicity is what is typically needed by the public health manager.

An alternative method that may satisfy the criteria listed above is the RBA. This competitive binding assay (Doucette et al. 1997; Ruberu et al. 2003) uses the same AOAC sample extraction procedure used for

the MBA. The 96-well plate format of the RBA allows testing of up to seven samples in triplicate, with three dilutions per sample to ensure the proper concentration range is represented. Multiple plates can be queued on the plate reader, with results from several successive plates available on the same day. In fairness, the MBA will likely provide results faster for the first several samples assayed, but will fail to meet the high throughput requirements during a major event due to the lack of automation. The RBA procedures are straightforward and can easily be performed by a trained technician. The reporting limit established in our laboratory for the RBA is significantly lower (4 µg/100 g tissue) than the detection limit of the MBA (35 µg/100 g tissue in the CDPH laboratory), illustrating the high sensitivity of the RBA method. Another advantage of the RBA is that it does not require careful filtration of samples prior to analysis as is the case with the HPLC method, reducing the time required for sample preparation. The majority of reagents are commonly available and relatively inexpensive, the exception being the tritiated STX needed for competitive binding. This reagent is not readily available through government services such as the National Institute of Standards and Technology (NIST), but is currently available commercially within the United States. Reliance on proprietary materials is always a point of concern for regulatory laboratories if there are no alternative sources available. A possible source of error in the RBA is the rat membrane synaptosome preparation. Not only is it a very inconvenient preparation procedure to carry out, but also due to its heterogeneity this membrane can be associated with high assay variability. One way to overcome this would be to have it available commercially as a standardised reagent.

Our previous experience with the RBA (Ruberu et al. 2003) was encouraging relative to the criteria mentioned above, and the precision of the method in our laboratory was found to be 10%. However, more recent work in our laboratory has suggested that method precision was no better than the MBA. Therefore, it was determined that a more detailed investigation into the various components of this assay was warranted in the hopes that method precision could be improved, facilitating the decision-making process for public health managers.

Materials and methods

Chemicals and reagents

- ³H-STX diacetate in methanol (Lot #040616, 0.1 mCi ml⁻¹, specific activity = 18.0 Ci mmol⁻¹) (American Radiolabeled Chemicals Inc., St. Louis, MO, USA).

- FDA reference standard, STX dihydrochloride (Lot #088 100 µg ml⁻¹ in 20% ethanol–water at pH 3.5) (USFDA, Office of Seafood, Laurel, MD, USA).
- Rat membrane synaptosome: the rat membrane preparation containing sodium channel receptors was composed of 20 brains from 6-week-old male Hotsman rats (Harlan Bioproducts, Indianapolis, IN, USA) and prepared according to the methodology of Doucette (Doucette et al. 1997). This preparation was divided into 2 ml aliquots and frozen at -70°C. A single aliquot was thawed for each RBA plate preparation.
- All reagents, standards and dilutions were prepared in 100 mM MOPS/100 mM choline Cl buffer at pH 7.4. To prepare this buffer, 20.9 g of MOPS (3-morpholinopropanesulfonic acid) and 13.96 g of choline chloride were dissolved in 900 ml of water, the pH adjusted to 7.4 and the final volume brought to 1 L with water.

Instrumentation

Scintillation counting was performed on a PerkinElmer Life and Analytical Sciences instruments TopCount® Model B. MicroScint-20 cocktail (PerkinElmer Life and Analytical Sciences, Shelton, CT, USA) was used as the scintillant for all RBA work.

Receptor binding assay (RBA) protocol

The RBA procedure involved the addition of 35 µl of MOPS/choline Cl buffer, 35 µl of unknown sample (or STX standard), 35 µl of ³H-STX, and 105 µl of a 1:6 diluted synaptosome preparation, in this order, to a 96-well microtitre filtration plate. A typical plate

outline is given in Figure 1. All calibration standards, QC samples, reference samples and shellfish sample extracts are run in triplicate on each plate. The first three columns of each plate were used to generate a calibration curve. Saxitoxin dihydrochloride standard was used for the calibration curve in the following final in-assay molar concentrations: 1 × 10⁻⁶, 1 × 10⁻⁷, 3 × 10⁻⁸, 1 × 10⁻⁸, 3 × 10⁻⁹, 1 × 10⁻⁹, 1 × 10⁻¹⁰ and 1 × 10⁻¹¹. Three wells per plate served as a reference blank, containing the material and reagents described above but omitting a source of non-radiolabelled STX. The reference blank establishes the maximum binding (*B*_{max}) for each plate. A quality control (QC) sample yielding an in-assay concentration of 3.0 × 10⁻⁹ M STX standard, independently made, was used as a daily QC check. All pipetting was carried out using a certified, calibrated eight-channel pipette. To achieve equilibrium binding, the plate was incubated for 1 h at 4°C, then filtered using a MultiScreen vacuum manifold system and rinsed with 200 µl of ice-cold (4°C) MOPS/choline Cl buffer to remove unbound toxin. To each well 50 µl of the scintillant (MicroScint®) were added, and the top of the plate sealed with tape. The prepared plate was placed inside the TopCount scintillation counter for 30 min. This allowed the scintillant to dark adapt and the contents to mix, prior to counting the receptor-bound ³H-STX.

Criteria that must be met for assay acceptance are as follows: (1) the slope of the standard curve must be between 0.8 and 1.2, (2) the relative standard deviation (RSD) of counts per minute (CPM) for each standard must be <30%, and (3) the QC check must be ±30% of the in-assay concentration of 3.0 × 10⁻⁹ M STX. Criteria for sample acceptance and quantification are: (1) *B*/*B*₀ = 0.3–0.7 and (2) RSD of the sample CPM must be <30%.

	1	2	3	4	5	6	7	8	9	10	11	12
A	1X10 ⁻⁶	1X10 ⁻⁶	1X10 ⁻⁶	Ref Blank	Ref Blank	Ref Blank	U3 1:10	U3 1:10	U3 1:10	U5 1:200	U5 1:200	U5 1:200
B	1X10 ⁻⁷	1X10 ⁻⁷	1X10 ⁻⁷	QC	QC	QC	U3 1:50	U3 1:50	U3 1:50	U6 1:10	U6 1:10	U6 1:10
C	3X10 ⁻⁸	3X10 ⁻⁸	3X10 ⁻⁸	U1 1:10	U1 1:10	U1 1:10	U3 1:200	U3 1:200	U3 1:200	U6 1:50	U6 1:50	U6 1:50
D	1X10 ⁻⁸	1X10 ⁻⁸	1X10 ⁻⁸	U1 1:50	U1 1:50	U1 1:50	U4 1:10	U4 1:10	U4 1:10	U6 1:200	U6 1:200	U6 1:200
E	3X10 ⁻⁹	3X10 ⁻⁹	3X10 ⁻⁹	U1 1:200	U1 1:200	U1 1:200	U4 1:50	U4 1:50	U4 1:50	U7 1:10	U7 1:10	U7 1:10
F	1X10 ⁻⁹	1X10 ⁻⁹	1X10 ⁻⁹	U2 1:10	U2 1:10	U2 1:10	U4 1:200	U4 1:200	U4 1:200	U7 1:50	U7 1:50	U7 1:50
G	1X10 ⁻¹⁰	1X10 ⁻¹⁰	1X10 ⁻¹⁰	U2 1:50	U2 1:50	U2 1:50	U5 1:10	U5 1:10	U5 1:10	U7 1:200	U7 1:200	U7 1:200
H	1X10 ⁻¹¹	1X10 ⁻¹¹	1X10 ⁻¹¹	U2 1:200	U2 1:200	U2 1:200	U5 1:50	U5 1:50	U5 1:50	QC	QC	QC

Figure 1. Layout of a typical 96-well plate used in RBA. The first three columns are used to generate the calibration curve. Six wells are used for quality control samples (QC) spiked at 3.0 × 10⁻⁹ M in assay concentration, three wells are used to determine maximum binding (ref blank) (*B*_{max}), and the rest of the wells are used for unknown samples (U).

Statistical analysis

MedCalc statistical software (MedCalc Software, Mariakerke, Belgium; Windows Version 10.4.8.0; <http://www.medcalc.org>) was used for all statistical analyses. Analysis of variance (ANOVA) was used to evaluate the significance of variability of mean counts among sequential plate readings and among detectors for a given plate reading. Plates found to have a significant difference among either sequential readings or detectors were subjected to post-hoc significance testing with the Student–Newman–Keuls (SNK) test for all pairwise comparisons. Prism (Graph Pad Software, Inc., La Jolla, California, USA) was used to generate the STX binding curves.

Results and discussion

Method variability study of RBA

Our laboratory has been following the RBA protocol as developed by Doucette et al. (1997) and later modified by Ruberu et al. (2003) for the detection of STXs in shellfish matrices. This methodology involves the competitive binding between STX analogues (in sample) and tritiated saxitoxin (^3H -STX) reagent. As we gained more experience with the assay it became clear that there were sources of variability that were not well understood. This involved unacceptable levels of variance among triplicate samples at a given dilution, variability in reference samples, and in QC standards placed at the beginning and end of each plate. The RBA requires pipetting of small volumes of reagents (35–100 μl) and is comprised of several independent steps, each of which is a potential source of variability. Our previous work (Ruberu et al. 2003) determined the RSD for assays of environmental samples to be 10%. In order to study the overall variability of the RBA with the goal of improving the method's precision, it was necessary to deconstruct the assay to its simplest components, then “rebuild” step by step, evaluating each step for its contribution to overall assay variability. Identified in this study are inherent differences among wells of the plate for replicate samples, heterogeneity of the rat membrane (binding sites) in each well, and the competitive binding process in each well. Given that each well acts as an independent experiment within a single plate, some amount of variability was expected for the measured CPMs among wells.

Another factor that can contribute to assay variability is the instrument's inherent variability among its 12 detectors, each of which reads a total of eight wells per plate. Detector normalisation is performed as part of routine maintenance of the instrument to minimise the variability that may exist among detectors. To understand detector variability it is important to know how the detectors are set up and which order the

detectors read the wells. The TopCount[®] instrument has two rows of six detectors each. The plate is read starting from the top row A to bottom row H (Figure 1). When a plate is read, the first set of six detectors measure wells A1, A3, A5, A7, A9 and A11, then move down to read wells B1, B3, B5, B7, B9 and B11. Subsequently, wells C1, C3, C5, C7, C9, C11 and A2, A4, A6, A8, A10 and A12 are read simultaneously by both sets of detectors. This continues until the set of wells G2, G4, G6, G8, G10 and G12 and the last set of wells H2, H4, H6, H8, H10 and H12 have been read by the second set of detectors. Not all wells are read simultaneously. As such, with a 5-min count time per well, the time difference between the measurement of the first and last wells is about 50 min. This can be a substantial period with respect to dissolution between sample and cocktail. To evaluate this potential source of variability to the assay, the count data for the series of plates studied were grouped by detector and statistically analysed by ANOVA to determine if there was a significant difference among the 12 detectors and, if so, which detectors were responsible for this variability.

Instrument background plate

To determine the inherent background variability in counts among the wells of a single plate, all 96 wells were filled with 50 μl of MicroScint[®] cocktail and counted three times in succession with a 30-min dark adapt delay period prior to each measurement. Background counts ranged from 8 to 36 CPM, from 7 to 27 CPM, and from 8 to 27 CPM for the three consecutive readings with average counts of 17.9, 17.8 and 16.5 CPM respectively. The standard deviation (SD) for the three count cycles ranged between 4 and 5 CPM. Figure 2 shows the CPM variability of the instrument background plate with respect to each

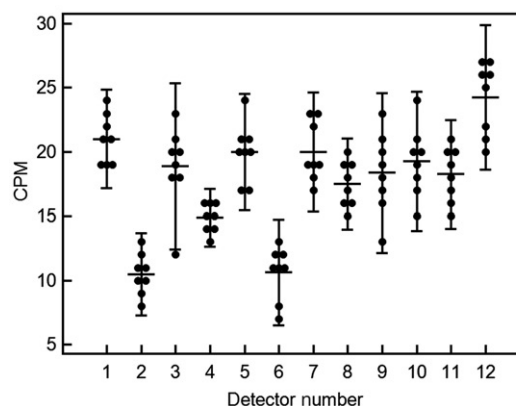


Figure 2. Plot of the instrument background plate second count cycle depicting the randomness of CPMs in the 96 wells. Each datum point represents the CPM of a well read by the respective detector. Also shown for each detector are the mean and error bars at 2 SDs for the group of data.

detector. Since there is no mixing of reagents involved in this plate, the variation seen here is attributed solely to counting statistics and to differences between the 12 detectors themselves. There was no significant pattern of variability observed for any single detector or to the time at which a well was counted. Although detectors 2 and 6 exhibited lower cpm values than the other ten detectors (Figure 2), the pattern of detector performance varied among the three sequential plate readings.

Blank plate

The next step was to determine the variability in counts among wells when a source of tritium was present. For this study all 96 wells were filled with 35 μl of ^3H -STX followed by 50 μl of MicroScint[®] cocktail. This blank plate was counted five times in succession with a 30-min dark adapt delay period prior to each counting cycle. Potential contributors to variability such as rat membrane preparation, competing non-labelled toxin, the competitive binding process itself and the washing/filtering step were absent.

Results showed a gradual increase in average CPM for the five sequential readings (Figure 3) with the greatest increase between the first ($\text{CPM}_{\text{average}} = 700$; $\text{RSD} = 19\%$) and second ($\text{CPM}_{\text{average}} = 869$; $\text{RSD} = 17\%$) measurements. The CPM stabilised with the next three readings ($\text{CPM}_{\text{average}} = 915, 939, 954$; $\text{RSD} = 17\%, 16\%, 16\%$). A one-way ANOVA determined that there was a significant difference among the mean CPMs ($p < 0.001$) for the five counting cycles. SNK post-hoc significance testing for all pairwise comparisons determined that the first two plate readings were significantly different from one another ($p < 0.05$) and both were significantly different from plate readings three through five. It also showed that the last three plate readings were not significantly different from one another ($p > 0.05$). From these

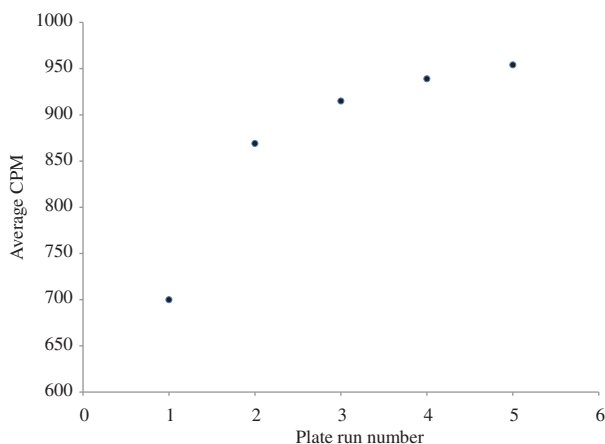


Figure 3. Graph of the blank plate five sequential readings showing a gradual increase in average CPM.

observations it is clear that the increase in CPM is due to mixing of the aqueous phase with the cocktail. An additional time of 3 h is needed to reach equilibrium in mixing. This is not a practical concern for the present assay because subsequent steps involve rinsing and filtering of each well prior to adding the cocktail, resulting in a single phase in each of the wells at the time of counting. However, shaking the RBA plate prior to the incubation step would make certain that all the reagents in the wells are properly mixed. These results give insight into the two-phase mixing process in a plate format. Unlike conventional liquid scintillation counting methods where 20 ml vials are vigorously shaken to obtain homogenous mixing prior to counting, in the plate format mixing can be an issue in obtaining reproducible results when assays with two phases are involved. This is further supported by looking at the first set of wells measured at the start of the count and the last set of wells measured (about 50 min later) within a single plate. The former (row A) had a $\text{CPM}_{\text{average}}$ of 447 while the latter (row H) had a $\text{CPM}_{\text{average}}$ of 723.

Reference plate (non-competitive binding)

The next step in reconstructing the assay involved introduction of binding sites for the ^3H -STX, i.e. non-competitive binding. In practice such a reference sample is run in triplicate on each RBA plate and the average CPM is used to determine maximum binding, B_{max} . This B_{max} value is used as the baseline and is compared with the sample CPM to generate the actual binding of samples. The reference plate was prepared by adding reagents in the following order: 35 μl of MOPS buffer, 35 μl of ^3H -STX and 105 μl of rat membrane preparation, then processed following the standard RBA protocol described above. This plate was measured three successive times.

The reference plate had a higher average CPM (1196 CPM) compared with the blank plate, with an RSD of 19%. Theoretically, the reference plate CPM values are expected to be lower than the blank plate, because the membrane binding sites would not retain all of the available ^3H -STX, with the excess being removed during the filtration step. The lower CPM of the blank plate is attributed to incomplete mixing of the ^3H -STX with the scintillation cocktail rather than the amount of tritiated toxin present. Since there is no aqueous phase in the reference plate, mixing does not become an issue. When the CPM values of the wells counted first (row A) are compared with those counted last (row H) there was no significant difference, which supports that phase mixing is absent. The comparable RSDs for the blank plate and the reference plate suggest that the addition of the rat membrane preparation, and the subsequent rinsing and filtering steps, do not contribute a significant amount of variability to

the assay. A one-way ANOVA determined that there was no significant difference among mean CPMs ($p > 0.3$) for the three sequential plate readings.

The reference plate, however, exhibited a significant difference among detectors (two-way ANOVA, $p < 0.001$). The same pattern in detector performance was observed for all three plate readings and the SNK pairwise comparisons determined that detector #12 was significantly different from all other detectors ($p < 0.001$) (Figure 4). The data from detector #12 were omitted and the statistical analysis repeated. The removal of this detector's data did not change the ANOVA outcome for sequential plate readings or detector variability.

QC plate (competitive binding)

To evaluate the added variance component associated with competitive binding, a non-labelled STX standard was added to compete with the $^3\text{H-STX}$, creating a competition for binding sites. For the non-labelled STX, a solution at 1.8×10^{-8} M, with a final concentration of 3.0×10^{-9} M in assay, was used. The standard RBA plate configuration contains triplicates of this solution and their average CPM is used as the plate's QC sample. The reagents added per well for the QC plate were identical to the reference plate described above, with the addition of 35 μl of QC sample prior to the addition of 35 μl of the $^3\text{H-STX}$. This plate was measured three successive times.

As expected, due to the introduction of competitive binding, the mean CPM of the QC plate was considerably lower than that of the reference plate (825 and 1196 CPM, respectively). Fewer binding sites for the radiolabelled toxin resulted in lower activity in the well after the rinsing and filtering steps. Triplicate counting of this plate gave an RSD of 17%. There appeared to be a slight decline in counts over the three successive

plate readings (Figure 5). There was a slightly significant difference among successive plate readings ($p = 0.04$), which was due to a significant difference between the first and third plate readings ($p < 0.05$).

Consistent with the results of the previous plate, there was a significant difference among detectors (two-way ANOVA, $p < 0.001$). The SNK pairwise comparisons of detectors did not identify a single detector to be different from all others, however detector #11 differed significantly from seven other detectors ($p < 0.05$) and detector #12 differed significantly from five other detectors ($p < 0.05$).

Overall assay variability

Introduction of the heterogeneous rat membrane preparation increased RSD only slightly for triplicate readings from 16% (blank plate) to 19% (reference plate). This demonstrates that the number of receptor sites in each aliquot of the membrane preparation is fairly uniform and does not affect assay precision significantly. With the introduction of competitive binding (QC plate) the RSD remained in the same range: 17%. Overall, an inherent variability of approximately 17% is associated with this assay, which is independent of the addition of the membrane preparation or the non-radiolabelled STX and subsequent competitive binding process. When assay variability was evaluated with respect to individual detectors, there was no reproducible pattern in detector performance, although there were frequent occurrences of one or more detectors having significantly different CPM than the rest of the detectors for a given plate. The detectors with the lowest and highest levels of variability were different from plate to plate. Removal of data for a detector that was found to be significantly different from a majority of the remaining detectors did not change the outcome of the ANOVA for any of the series of plates. This detector variability observed

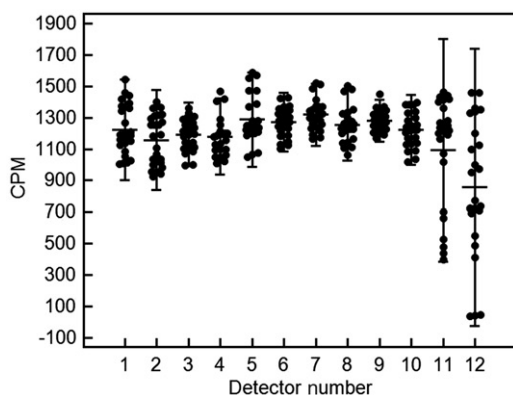


Figure 4. Scatter plot of the reference plate, first reading showing detector variability. Each datum point represents the CPM of a well read by the respective detector. Also shown for each detector are the mean and error bars at 2 SDs for the group of data.

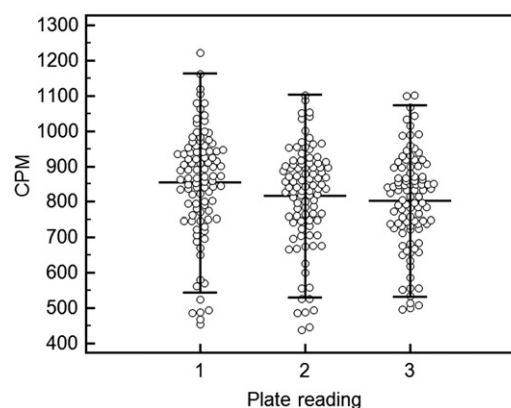


Figure 5. Results of the QC plate showing the variability of CPM in the three successive readings. For each run the mean and error bars at 2 SDs per plate are shown.

could be attributed to pipetting error along single rows. Since an eight-channel micropipette is used to add reagents to single rows, an error in one such addition will show up as a single detector inconsistency. This would erroneously label one or more detectors as being different to other detectors. Although our initial work (Ruberu et al. 2003) on the RBA showed an RSD of 10%; a more recent study (van Dolah et al. 2009) demonstrated an RSD of 17.7% comparable with the current finding of 17%. It should be noted that our initial work was conducted using a six-detector instrument and with more experienced analysts, which could be the reason for the lower RSD for that study.

RBA acceptance criteria

In a typical RBA plate, each sample (calibration standard sample, QC sample, unknown sample, reference sample) is run in triplicate and the average CPM value is used for further calculations. Triplicate samples, rather than duplicates, are run to improve the accuracy of this assay. According to the acceptance criteria of the RBA assay (van Dolah et al. 2012), a given set of triplicate sample data are rejected if the RSD exceeds 30%, requiring reanalysis of the rejected sample. On average about 10% of the samples analysed in our laboratory are rejected due to the high variance among the triplicate values. In addition, if the QC sample CPM has >30% RSD, then the entire plate must be rejected. This loss of data results in the need to prepare and run a new plate, increasing both the turnaround time for results and the cost of the assay. One possible way of preventing samples from being rejected is to identify and remove outliers within a set of replicates. By eliminating outliers, the variability of replicates may be reduced to an acceptable level (<30% RSD), preventing invalidation of the entire plate or of individual samples. Therefore, we investigated a statistical approach to eliminate outliers methodically.

Grubbs' test and Student's t-test

A comparison of statistical outlier tests concluded that the Grubbs' test (Grubbs 1969) and the Student's *t*-test

(Sokal and Rolf 1981) were best suited for determining an outlier within a triplicate dataset. The Grubbs' test compares the suspected outlier to the mean of all replicates, including the suspected value. The Student's *t*-test compares the potential outlier to the mean of the remaining values. The Grubbs' test is therefore more conservative in approach and it would be expected that this test would identify fewer outliers than the Student's *t*-test. The Grubbs' test for triplicates determines that a value is an outlier if the calculated value (*G*) is greater than the critical value (*Z*) of 1.153 at a 95% confidence interval ($\alpha = 0.05$). The Student's *t*-test determines that a value is an outlier in a triplicate dataset if the calculated *t*-value is greater than the critical *t*-value of 12.706 ($\alpha = 0.05$). By running the triplicate sample data values through these statistical tests, an outlier can be determined in an unbiased fashion, possibly avoiding the rejection of the entire sample.

The two outlier tests were evaluated by analysing data of each plate reading for the experiments presented above. For example, statistical analysis of the second plate reading of the QC plate, which had an RSD of 17.5% with an average CPM value of 817, resulted in both tests identifying a total of four outliers (Table 1). The results calculated after the four outliers were removed gave an average of 823 CPM with a 17% RSD. Removing the outliers did not improve the RSD significantly but showed a slight increase in CPM. Other plates tested for outliers had similar results, with an insignificant lowering of the sample RSD and slight increases or decreases in CPM. Since the CPM values directly relate to STX concentration of a sample, it is possible that the removal of outliers could have an impact on the accuracy of the assay results.

In order to evaluate how outlier testing would impact results of actual shellfish samples, 17 samples were spiked with STX concentrations that ranged from 5 to 1000 µg/100 g shellfish tissue and were evaluated for outliers using both statistical tests. Each triplicate set of data were assessed in two ways: raw data (no outliers removed) and data with outliers removed. Each test identified the same outliers and, as expected, the *t*-test identified additional outliers that were not detected by the Grubbs' test. Removal of the outliers kept these samples from being rejected (<30% RSD in

Table 1. Results from two outlier tests for three successive readings of the QC plate.

Plate reading	Raw data			Grubbs test outliers removed			Student's <i>t</i> -test outliers removed			RSD % change	
	Mean CPM	SD	RSD	Mean CPM	SD	RSD	Mean CPM	SD	RSD	Grubbs	<i>t</i> -test
1	854	155	18	859	152	18	859	152	18	0.03	0.03
2	817	143	18	823	140	17	823	140	17	0.03	0.03
3	803	135	17	810	135	17	809	136	17	0.01	0.00

triplicate wells), however there was no significant improvement in the precision of the assay. The average recovery of STX was 118% and 113% for raw data and data with outliers removed, respectively. Although the assay accuracy was improved on average, some plates showed a decrease in accuracy after outliers were removed, indicating that the suspected outlier was closer to the actual value than the remaining data points. In practice, outlier testing would likely reduce the number of samples and plates rejected, thereby reducing the time required to report results and lowering the per sample cost of the assay. The potential negative effect on method accuracy suggests caution with this approach in the absence of tangible evidence of analytical error during plate preparation.

Control charts for RBA

A better way of identifying erroneous data is through the use of control charts, which are based on a laboratory's acceptable and attainable performance criteria for precision and accuracy for a given method. A control chart enables the laboratory to monitor its performance visually by updating the chart with data from each subsequent analytical run. In this way a control chart for each critical parameter of a method tracks the detection of data outside of the acceptable performance limits. Control charts are prepared by plotting the date or run number as the abscissa and the value of interest, e.g. STX concentration of the QC check sample estimated on each plate, as the ordinate. Performance limits are established by averaging at least 20 measurements that have acceptable individual statistics, setting control limits and identifying the range of variability for that parameter. Rather than setting an arbitrary acceptance limit of $\pm 30\%$ for recovery of the QC sample, each laboratory can establish control limits based on their performance to determine whether or not an RBA plate is acceptable. Typical control limits are based on the number of SDs from the estimated mean. Once the mean and SD have been determined, the parameters from each subsequent assay are added to the appropriate control chart to maintain a continuous record of performance. In addition to the detection of erroneous values that would indicate an unacceptable plate, control charts allow tracking of systematic changes in method performance (e.g. due to degrading stock solutions, changes in materials like plate manufacturers, etc.) as well.

Figure 6 shows control charts for three RBA parameters acquired from our laboratory over a period of 1 year: (1) QC check standard, (2) slope of the binding curve and (3) EC_{50} . For each of the three control charts, control limits were based on the mean ± 2 SD of the first set of 20 acceptable data. For example, the estimated mean for the QC check standard (3.0 nM theoretical concentration) from the

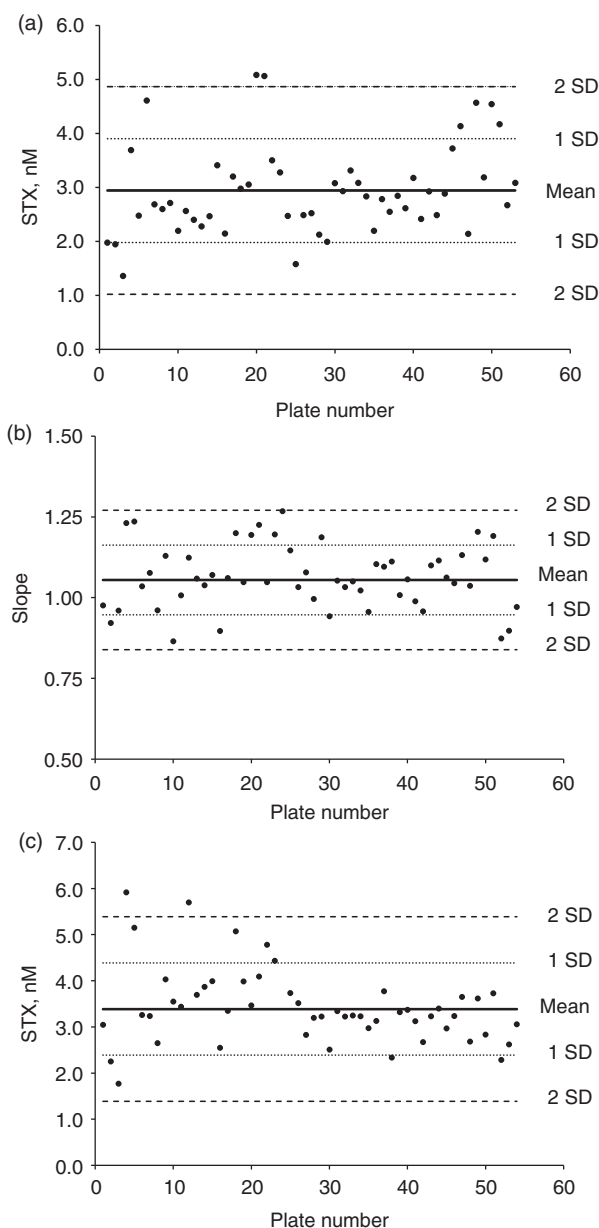


Figure 6. Control charts for (a) averaged daily QC samples on a plate, (b) slope of the binding curve and (c) EC_{50} per plate.

first 20 plates was 2.94 nM and the calculated SD was 0.962 nM, resulting in control limits of between 1 and 4.5 nM (Figure 6(a)). Two data points on this plot had QC check standard values that were found to be outside of the control limits, requiring those two plates to be rejected and the assay run again for those samples. In contrast, the current RBA protocol of $\pm 30\%$ calculates an acceptance limit of between 2.1 and 3.9 nM for the QC check standard, which would result in 12 data points out of control, hence the rejection of 12 RBA plates. The latter criterion is arbitrary since the accuracy of the QC sample is method, instrument and analyst specific and must be established per individual laboratory.

When the established control limits are exceeded and a trend is observed, results are investigated for method bias and potential mistakes, allowing corrective actions to be taken to address the root cause to prevent recurrence of the error. Figure 6(b) shows the control chart for the slope of the standard curve. The slope was demonstrated to be a very stable parameter, with tight control limits ranging from 0.8 to 1.3 with no data points being rejected. The EC_{50} parameter is shown in Figure 6(c). In this case a high variability is seen at the beginning of the chart and as the analyst gains more experience with the assay the EC_{50} value becomes more consistent with less variation.

An additional parameter that has been inconsistent and highly variable in the RBA is the maximum binding, or top value of the binding curve. Ideally, the standard binding curve should plateau at 100% binding. However, often times we observe the plateau significantly below (80%) or above (120%) this value. Such a large shift in the curve significantly affects the final results of STX concentration in a sample. It is sometimes caused by one or more of the lowest three standards being out of control, thus dragging the curve in one direction. The top plateau was monitored using a control chart (Figure 7). The 1 SD control limit established for this data was between 0.9 and 1.1, which is $\pm 10\%$ binding. We have observed that this parameter can have a significant effect on the outcome of the results and therefore recommend developing a control chart to monitor its performance. Currently the importance of the top plateau is overlooked and not considered as an assay performance acceptance criterion.

Conclusions

Previous studies from our laboratory demonstrated the RBA method for the detection of STXs to be very reliable and to have the potential of being an alternate

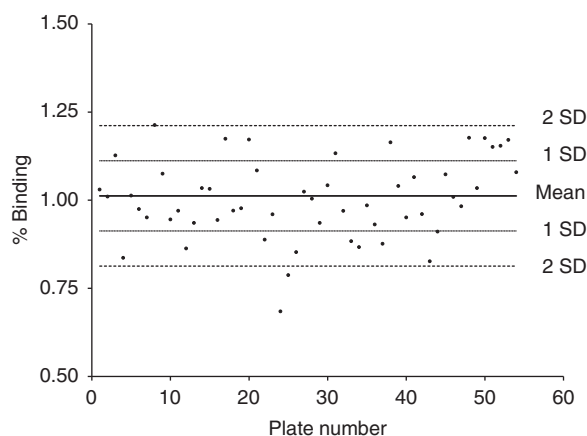


Figure 7. Control chart for the top plateau, maximum binding (B_{max}), of the calibration curve.

regulatory test method for PSP. Our current work focused on identifying sources of variability associated with the RBA and evaluating alternate QC approaches for validating test plates. The assay variability work included evaluation of each step of the assay by deconstructing its procedural steps, and also assessing the instrument's detector variability. The overall variability of the assay was determined to be 17%. Results discussed above show that the variability within a plate arises from several factors, such as counting statistics, analyst variability, mixing of well contents with cocktail, and the inherent measurement technique of the TopCount[®]. It is not known if the same variability would be observed in other instruments with different numbers of detectors or with detectors placed in a different array. A pipetting error along a row by the eight-channel micropipette would point to single detector variability as well and would be hard to identify. It is recommended periodically to evaluate individual detector performance with either a reference plate or a QC plate format similar to that used in the current study.

We have explored the use of two different outlier tests, Grubbs' test and Student's *t*-test, alone and in combination with the allowable procedure recommended in the NOAA protocol. Overall, removal of outliers lowers the RSD between replicate wells of a sample to $<30\%$, thus preventing that sample from being rejected. As a result it is expected that routine outlier testing would reduce the number of samples and plates rejected under the current QC criteria, which would help minimise the turnaround time between sample receipt and the reporting of results. The reduction in the number of rejected samples would also lower the cost per sample of the assay. Although some improvement in precision will be gained when an outlier is excluded, it is possible that accuracy will be diminished if the excluded value is closer to the actual target concentration. The potential negative effect on method accuracy suggests that the removal of suspected outliers should only be considered if it is suspected that there is an error associated with the sample(s) involved (e.g., a pipetting error).

Another avenue explored was the use of control charts for monitoring the three critical parameters of the RBA method, i.e. QC check standard, slope and EC_{50} . Establishing acceptable limits within each laboratory for respective parameters will ensure consistent performance over time, identify plates that must be rejected because one or more parameters are outside of the set control limits, and allow identifying and correcting process changes that would affect every assay. Establishing control limits for the maximum binding (B_{max}) as a fourth critical parameter for RBA performance is recommended. Such a development of associated control charts can be a part of the

laboratory's routine QC programme and is recommended as the primary quality control process for the RBA.

Acknowledgements

This study was supported under NOAA grant NA04NOS4780239 from the Monitoring and Event Response for Harmful Algal Bloom (MERHAB) programme. This is MERHAB Publication Number 152. The authors would like to thank Roger Ho for helping with the Graphpad Prism data analysis; Clive Kittredge and Vanessa Zubkousky for the preparation and analysis of plates; and American Radiolabeled Chemicals Inc. for providing ³H-STX reagents. Special thanks to Sherwood Hall of the US Food and Drug Administration's Office of Regulatory Science for FDA STX reference standards and helpful discussions on this study. The authors would also like to thank Fran van Dolah and Greg Doucette for their guidance in the RBA method.

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Single Laboratory Validation (SLV) Submission to the Interstate Shellfish Sanitation Conference (ISSC) in support of Method Approval as an Approved NSSP Method

Justification for New Method

For: Domoic Acid (ASP) Plate Kit, Cat. # 20-0249

Type of Method: Enzyme Linked Immunosorbent Assay (ELISA) utilizing a polyclonal antibody for detection of the ASP Biotoxin, Domoic Acid.

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Date of submission: June 30, 2017

Purpose of Intended Use of the Method. The purpose and intended use of this method is to provide a laboratory method for quickly establishing a quantified level of the ASP biotoxin, Domoic Acid, in *Mytilus edulis* (Blue Mussel) tissue as required for closing and opening of shellfish growing areas.

Need for the New Method in the NSSP, Noting Any Relationships to Existing Methods. It appears there is a need for additional approved methods as an alternative to HPLC for the Biotoxin type: Amnesic Shellfish Poisoning (ASP). An ELISA method would enable monitoring laboratories to become proficient in operating a quantification method for ASP toxins levels at a lower equipment and reagent cost using a method that requires less training than HPLC to operate. This proposal demonstrates the equivalency of the HPLC and ELISA when using the same sample extract. This offers an additional benefit in that any confirmation testing could be completed using the same sample extract.

Method Limitations: This proposal offers supporting data for use of the method with *Mytilus edulis* (blue mussel) tissue only.

Method Documentation

Method Title: Domoic Acid (ASP) Plate Kit, Cat. # 20-0249

Method Scope: The method is a competitive enzyme linked immunosorbent assay (ELISA) for the quantification of domoic acid (DA) residues in *Mytilus edulis* shellfish tissue. Domoic acid is produced by some species of the diatom *Pseudo-nitzschia* which is the primary toxin associated with amnesic shellfish poisoning (ASP). Current legislation in the NSSP limits the amount of DA allowed in harvested shellfish to 2 mg/100 g (20 ppm) and will close shellfish growing areas to shellfish harvesting to protect consumers from exposure to the toxin. The test kit provides a tool to close and open shellfish growing areas by rapidly monitoring toxin levels as levels can quickly rise and fall.



References: (For HPLC Method) M.A. Quilliam, M.Xie and W.R. Hardstaff. 1991. Rapid Extraction and Cleanup Procedure for the Determination of Domoic Acid in Tissue Sample. NRC Institute for Marine Biosciences, Technical Report #64 National Research Council Canada #33001.

ELISA Principle- The kit is a competitive ELISA method utilizing a plate coated with Rabbit polyclonal antibodies specific to the analyte Domoic Acid. The Domoic Acid–HRP (Horse Radish Peroxidase) Enzyme Conjugate competes with any toxin from the shellfish sample extract for binding with the antibody on the plate. After an incubation period of 30 min, the plate is washed with water to remove excess material and an enzyme substrate is added to react with any HRP Enzyme Conjugate bound to the plate. The amount of bound Conjugate is inversely proportional to the amount of DA toxin in the sample extract and can be visualized by development of a blue color. The ELISA is stopped at 30 min. with a 0.1 N HCl solution, and evaluated by reading the absorbance (OD) at 450 nm wavelength in a plate reader. The OD of the sample is compared to the Calibration Curve and multiplied by the total dilution factor of 4000 to obtain the concentration of toxin in the shellfish tissue sample.

Shellfish Sample Preparation: Fresh shellfish are externally washed and removed from the shell, approximately 15 single animals are combined. Composite sample is washed, drained then homogenized for ~20 seconds using a Waring blender with 16 oz. Mason jar fitted with ice crusher blade. Samples are aliquoted and can be frozen at -20°C until use.

Shellfish Sample Extract Preparation: Composite mussel samples are extracted using a 4 X ratio of 50% methanol/water to tissue weight and mixed using a vortex mixer for 3 minutes. A sample of approximately 1 ml was aliquoted into a 1.5 ml Eppendorf tube and centrifuged for 5 min. at 12,000 rcf. Supernatants were diluted 1:1000 (as directed in the Test Kit Product Insert) into 10% acetonitrile/water (Sample Dilution Buffer). Diluted samples are used in the ELISA. The extraction and dilution procedure results in a total dilution factor of 4000 to be used in calculation of DA residues present in the original tissue sample.

Proprietary Aspects. Beacon Analytical Systems has developed the kit including antibodies and HRP enzyme conjugate.

Equipment: Microplate Reader with a filter for reading at 450 nm wavelength. Sample Preparation- blender, scale, extraction container with lid (10-20 ml), vortex mixer, microcentrifuge (12,000 rcf), disposable 1.5 ml centrifuge tubes, calibrated variable pipettes 1.0 ml and 0.010 ml with disposable tips, timer, and wash bottle.

ELISA Kit Reagents.

- **Plate** – (1) containing 12 test strips of 8 wells each vacuum-packed in aluminized pouch with indicating desiccant.
- **Domoic Acid Calibrators**– (4) vials each containing 2 ml with a concentration of 0, 0.5, 5, and 50 µg/L (ppb) Domoic Acid
- **Domoic Acid HRP Enzyme Conjugate** – (1) vial containing 12 ml
- **Substrate** – (1) vial containing 14 ml
- **Stop Solution** – (1) vial containing 14 ml (Caution! Contains 1N HCl. Handle with care.)
- **Product Insert** containing instructions for use.
- **Certificate of Conformity** (Specific to each Kit Lot#).



ELISA Test Procedures:

1. Allow reagents and sample extracts to reach RT prior to running the test
2. Place the appropriate number of test wells into a micro well holder. Be sure to re-seal unused wells in the zip-lock bag with desiccant.
3. Using a pipette with disposable tips, dispense **100 µl** of the **Calibrator or sample extract** into the appropriate test wells. Please use a clean pipette tip for each sample addition.
4. Dispense **100 µl** of the **HRP Enzyme Conjugate** into each well.
5. Shake the plate gently for 30 seconds using a back and forth motion. Then incubate the wells for **30 minutes** at RT.
6. Decant the contents of the wells into an appropriate waste container. Fill the wells to overflowing with laboratory grade water and then decant. Repeat four times for a total of five washes.
7. Following the last wash, tap the inverted wells onto absorbent paper to remove the last of the water.
8. Dispense **100 µl** of the **Substrate** into each well. Shake the plate gently for 30 seconds using a back and forth motion.
9. Incubate the wells for **30 minutes** at RT.
10. Dispense **100 µl of the Stop Solution** into each well.
11. Measure and record the absorbance (Optical Density; OD) of the wells at 450 nm using a strip or plate reader. The OD correlates to a concentration of DA (ppb) based upon the Calibration Curve run with each set of samples.
12. To obtain the concentration of Domoic acid in the sample multiply the concentration results by the Total Dilution Factor of 4000.

Note: If the sample absorbance is higher or lower than the 0.5 or 50 ppb Calibrator results, the tissue levels should be expressed as less than or greater than the corresponding tissue levels (<2ppm or >200ppm DA). The sample dilution can be modified appropriately and retested along with another set of Calibrators.

Note: Running Calibrators and samples in duplicate will provide optimal assay precision and accuracy.

Quality Control:

Beacon Analytical Systems is ISO 9001:2015 approved for their quality systems for immunochemical test kit development, manufacturing and supporting activities.

Overview of Kit Quality Control - Each kit is tested following the ELISA procedure in the product insert. During manufacturing operation duplicates of the Kit Calibrator Reagents are run in order to meet established criteria prior to shipment.

Each Domoic Acid (DA) Calibrator's absorbance (OD) and binding characteristics (% B/B0) must be within a set of ranges. Ranges have been established for the Domoic Acid Plate Kit from historical data.

All plate components are tested for precision prior to using them in kits. The tolerance for variation within one lot of plates is less than or equal to 5%.

DA Calibration solutions are prepared using certified reference standard material purchased from the Canadian National Resource Council and are tested to be within 2% of the previous lot of control.

The R² correlation of the DA Kit Calibration Curve should be 0.99 or above.

All CV's must be less than or equal to 5%.

All QC data is kept electronically and backed up with hard copies at our manufacturing plant.



Single Laboratory Validation Criteria and Results

Section # 1 - Accuracy/ Trueness & Measurement Uncertainty

Section # 2 – Ruggedness

Section # 3 - Precision & Recovery

Section # 4 - Specificity

Section # 5 - Linear Ranges, Limit of Detection, Limit of Quantitation / Sensitivity,

Section # 6 - Comparability (If intended as a substitute for an established method accepted by the NSSP).

Materials and Methods

Blank Mussel Tissue: Five different bags of mussels were purchased locally and screened on the ELISA for DA content. The ELISA screening did not find DA residues in these samples. They were used in spiking experiments (Table 1).

Table 1: Sample Type - Mussel (Blank) DA Blank Mussel Tissue used in validation.

ID	Harvest Date	Location	Type	Commercial Name	DA ELISA Screening Results*
A.	10/24/2016	Chebeague Island, ME	Aquaculture	Bangs Island Mussels	0 ppm
58	01/30/2017	Addison, ME	Natural	Moosabec Mussels	0 ppm
59	02/02/2017	Chebeague Island, ME	Aquaculture	Bangs Island Mussels	0 ppm
F.	04/18/2017	P.E.I., Canada	Natural	Cape Cod Shellfish Co.	0 ppm
E.	04/24/2017	Addison, ME	Natural	Moosabec Mussels	0 ppm

*All Blank tissue screening results were below the DA ELISA Level of Quantitation (LOQ).

Domoic Acid Standards

Certified Reference Material (CRM) – Certified Calibration Solution for Domoic Acid (CRM-DA-g, Lot# 20140730) purchased from National Research Council Canada. The certified concentration values and associated uncertainties for Domoic Acid and *epi*-Domoic Acid in this solution is 103.3 ± 3.4 ug/ml (at 20 °C). This is used in the manufacturing of Calibrator Solutions supplied with test kit.

Domoic Acid (DA) Standard used for sample spiking at ppm levels - Domoic Acid from *Mytilus edulis*, Calbiochem Catalog # 324378, Lot #2879693, 1 mg/vial. Reconstituted Solution adjusted for purity (98%).

Equivalency of CRM and DA Standards using HPLC Analysis: HPLC was used to confirm the concentration and equivalency of this standard to the reference material prior to the use in spiking experiments. The Calbiochem standard was diluted 1:40 into 10% acetonitrile/DI water (ELISA sample dilution buffer) or 50% methanol/50% DI water (Sample extraction solution) and each run in triplicate on HPLC. The concentration of the Calbiochem solution used for tissue spiking was assigned using the average of 6 replicates as 1.873 mg/ml.

Shellfish Sample Extract Preparation: Composite mussel samples are extracted using a 4 X ratio of 50% methanol/water to tissue weight. Samples were spiked with Calbiochem DA Standard at this time (if



required) and mixed using a vortex mixer for 3 minutes. A sample of approximately 1 ml was aliquoted into a 1.5 ml Eppendorf tube and centrifuged for 5 min. at 12,000 rcf. Supernatants were diluted 1:1000 (as directed in the Test Kit Product Insert) into 10% acetonitrile/water (Sample Dilution Buffer). Diluted samples were used in the ELISA resulting in a total dilution factor from the sample preparation of 4000.

Mussel Tissue- Certified Reference Material for Domoic Acid in Mussel tissue. - Certified Reference Material for Domoic Acid (CRM-ASP-MUS-d, Lot# 201112) purchased from National Research Council Canada. The concentration of DA and epi-DA is reported to be 49 ± 3 ug/g.

Extraction efficiency using Mussel Tissue CRM. Evaluation of Test method extraction and recovery was evaluated using this reference material (Table 2). Sample A was prepared by addition of CRM Reference mussel tissue to blank mussel tissue (Table 1) at a 1:1 ratio resulting in a DA tissue concentration of 24.5 ug/g, Sample F was prepared by blending the 4 gm standard with 9 gm of blank mussel tissue to obtain a tissue concentration of 15 ug/g. Both samples were extracted as described above and tested by ELISA. Recovery percentages at two different spiking levels were found to be 107 and 109 percent in mussel tissue by ELISA.

Table 2. – Mussel CRM extraction recovery results.

Sample	DA ELISA Result (ppm)	% Recovery
Blank Mussel Spiked with DA Mussel CRM		
Sample A spiked at 24.5 ppm DA	26.2 ppm	107 %
Sample F spiked at 15.0 ppm DA	16.3 ppm	109 %



Section 1: Accuracy/Trueness & Measurement Uncertainty (Table 3)

Working Range – Twenty samples of DA Blank Mussel Tissue were spiked with a low level (10 ppm), and twenty samples at a high level-20 ppm using the Calbiochem standard and extracted and evaluated by ELISA. Data and results are shown in Table 3.

Data Summary- Accuracy/Trueness

% Accuracy 10 ppm spike = 96.0 %

% Accuracy 20 ppm spike = 95.9 %

Data Summary – Measurement Uncertainty

Measurement uncertainty determined using a two-sided, 95% Confidence interval calculation

10 ppm spike = 0.662

20 ppm spike = 1.224

Table 3. Results of Accuracy/Trueness Testing of Blank and Spiked Mussel Tissue

Sample	Spiked Mussel (10 ppm)	% Accuracy (10 ppm)	Sample	Spiked Mussel (20 ppm)	% Accuracy (20 ppm)
1	10.70	107.0	1	21.06	105.3
2	8.06	80.6	2	23.97	119.8
3	8.07	80.7	3	16.87	84.3
4	10.29	102.9	4	19.07	95.3
5	11.15	111.5	5	16.27	81.4
6	8.18	81.8	6	17.18	85.9
7	8.43	84.3	7	16.80	84.0
8	11.26	112.6	8	18.62	93.1
9	11.42	114.2	9	16.31	81.5
10	8.81	88.1	10	22.74	113.7
11	8.91	89.1	11	20.10	100.5
12	9.51	95.1	12	18.06	90.3
13	97.7	97.7	13	21.85	109.3
14	10.60	106.0	14	17.25	86.2
15	8.63	86.3	15	17.55	87.8
16	12.20	122.0	16	22.39	111.9
17	9.42	94.2	17	19.16	95.8
18	8.39	83.9	18	17.06	85.3
19	8.02	80.2	19	22.27	111.4
20	10.18	101.8	20	18.96	94.8
Average	9.60	96.0 %		19.18	95.9 %
+/- SD	1.31			2.42	
Measurement Uncertainty @ 95% CI		0.662		Measurement Uncertainty @ 95% CI	1.224



Section 2: Ruggedness

Method: Composite mussel samples were spiked at 10 and 20 ppm, extracted with 50% methanol/water for 3 minutes, centrifuged and diluted in 10% acetonitrile/water with a total dilution factor of 4000. The diluted sample extract was evaluated on two different manufactured test kit lots. The data and results can be found in Table 4.

Data Summary:

Value for the test of symmetry of the distribution of Kit Lot 1	0.153
Value for the test of symmetry of the distribution of Kit Lot 2	0.563
Variance of kit Lot 1	26.07
Variance of kit Lot 2	49.53
Ratio of the larger to smaller variance of Lot 1 & Lot 2	1.89
Significant Difference between Lot 1 & Lot 2 based upon paired t-test	No

Table 4

Time of Analysis	Sample	Kit Lot 1	Kit Lot 2
Day 1	1A	8.43	8.81
	1B	6.94	8.91
	2A	11.26	9.51
	2B	11.42	9.77
Day 2	3A	9.51	9.25
	3B	10.6	9.42
	4A	8.63	8.39
	4B	12.2	7.86
Day 3	5A	10.48	8.77
	5B	10.18	9.99
Day 1	6A	17.18	22.74
	6B	16.8	29.36
	7A	18.62	23.97
	7B	16.31	25.41
Day 2	8A	22.39	16.87
	8B	19.16	19.07
Day 3	9A	21.06	16.27
	9B	17.23	24.37
	10A	20.77	17.4
	10B	22.27	19.1
Skewness		0.153	0.563



Variance	26.07	49.53
Ratio of variances	1.89	
P-Value (Paired t-test)	0.546	
Significant Difference	No	

Results: The data summary indicates the values of symmetry for kit lot 1 and 2 are within the range of -2 to +2, a non-significant degree of skewness in the distribution. The ratio of the variances between lot 1 & 2 is less than 2 indicating homogeneity of variance. A paired t-test used for data analysis results in a p-value of 0.546 which indicates there is no significant difference between Kit 1 and Kit 2.

Ruggedness continued - ELISA parameters

Method: Composite mussel samples were spiked at 10 and 20 ppm, extracted with 50% methanol/water for 3 minutes, centrifuged and diluted in 10% acetonitrile/water with a total dilution factor of 4000. The ELISA Standard Operating Procedure (SOP) parameters were modified in the ELISA then tested for an evaluation of the critical steps in procedure (Table 5).

1. Incubation time for the initial step of the ELISA is set at 30 min. The incubation time was modified to be a total of 15 minutes or 45 minutes. The spiked sample data was evaluated by Welsh’s t-test and found not to be significant at either time point tested.
2. A wash step is required in the ELISA to remove unbound materials. The SOP wash is repeated 5 times with water. This was changed to a 4 time wash. The spiked sample data was evaluated by Welsh’s t-test and found to be significant from the SOP data.
3. The SOP for test incubation temperature is that the ELISA should be run at RT (20-28 °C). The incubation temperature was modified to be 4 °C or 30 °C. The spiked sample data was evaluated by a paired t-test and found not to be significant at either temperature tested.
4. The kit reagents should be equilibrated to RT prior to running the ELISA. All kit reagents were removed directly from a 4 °C refrigerator and run in comparison to RT reagents. A t-test on the resulting data indicated no significant difference in the results.

Table 5

	ELISA Standard Operating Procedure	Definition of ELISA SOP	Variation Factor	Significantly different to SOP by t-test.	Variation Factor	Significantly different to SOP by t-test.
1.	Primary Incubation Time is 30 min.	Incubation time for HRP Enzyme Conjugate, Sample Extract or Calibrator on plate	Incubation time changed to 15 min.	No	Incubation time changed to 45 min.	No
2.	Plate water Wash Step is Repeated 5 times.	Water wash step to remove unbound materials prior to	Wash Step is changed to repeat 4 times.	Yes		



		Substrate addition				
3.	Incubation Temperature done at room temperature	ELISA incubation steps run at RT (20-28 °C).	ELISA Incubation at 4 °C.	No		ELISA Incubation at 30 °C.
4.	Reagent Temperature	Kit reagents are equilibrated to RT prior to running test.	Reagent Temperature is cold (4 °C).	No		

Section 3: Precision & Recovery

Precision

Method: Evaluation of mussel tissue spiked with a low (10 ppm), medium (20ppm) and high level (40 ppm) of DA was completed using the method of extract preparation and ELISA analysis previously outlined, to evaluate the method consistency over a range of concentrations.

Data Summary- Precision -The F value obtained in the evaluation between groups was less than the critical value of 2.39 (for 9 and 20 degrees of freedom) at 0.05 significance level indicating the mean values from the samples are not significantly different.

The F value obtained in the evaluation of different concentrations (subgroups within groups) is greater than the critical value of 1.93 (for 20 and 30 degrees of freedom) at the 0.05 significance level indicating the mean values of each concentration are significantly different. This is an expected result since there were three sample concentration used to generate the data (10, 20 and 40 ppm) which are quite different.

From this ANOVA analysis (Table 6) we can conclude that the precision of the method is consistent over the range of sample concentrations tested.

Table 6. **Fully nested/hierarchical random analysis of variance (ANOVA)**

<u>Source of Variation</u>	<u>Sum Squares</u>	<u>DF</u>	<u>Mean Square</u>
Between Groups	30.651802	9	3.405756
Between Subgroups within Groups	9,583.973276	20	479.198664
Residual	323.251852	30	10.775062
Total	9,937.87693	59	

F (VR between groups) = 0.316078 P = 0.9633

F (using group/subgroup msqr) = 0.007107 P > 0.9999

F (VR between subgroups within groups) = 44.472939 P < 0.0001



Recovery

Method: Evaluation of mussel tissue spiked with a low, medium and high level of DA was done using the method outlined, to evaluate the method consistency over a range of concentrations. The results are found in Table 7.

Data summary

The variance ratio for the component of concentration in sample is not significant at 95% CI.

Recovery Percentage over the average data set (10, 20 & 40 ppm) using spiked mussel tissue is 99.55%.

Table 7 RECOVERY

Sample	Low Spike 10 ppm		Medium Spike 20 ppm		High Spike 40 ppm	
	Average	Spike minus Average	Average	Spike minus Average	Average	Spike minus Average
1	8.62	1.38	19.96	0.04	37.00	3.00
2	7.93	2.08	23.08	-3.08	39.42	0.58
3	10.39	-0.39	21.30	-1.30	40.02	-0.02
4	10.60	-0.59	20.86	-0.86	37.39	2.61
5	9.38	0.62	19.63	0.37	39.00	1.00
6	10.01	-0.01	19.12	0.88	40.41	-0.41
7	8.51	1.49	18.67	1.34	43.41	-3.41
8	10.03	-0.03	20.80	-0.80	36.52	3.48
9	9.63	0.38	19.09	0.91	43.80	-3.80
10	10.09	-0.09	20.69	-0.69	41.52	-1.52

Anova: Single Factor

Source of variation	df	SS	MS	F	P-value	F crit
Concentration	2	3.24	1.62	0.55	0.59	3.35
Error	27	80.24	2.97			
Total	29	83.48				

Section 4: Specificity

Method: Four compounds were evaluated to challenge the specificity of the ELISA, three were included due to their similarity of structure to DA, glutamine & glutamic acid at 100 ppm and kainic acid at 20 ppm. Saxitoxin (20 ppm) was evaluated due to the potential that it may be present in the shellfish at the same time as DA. Shellfish extracts containing DA from extracted tissue levels of 0 (blank), 10 and 20 ppm DA were run on the ELISA. These same extracts were run in the presence of the suspected interfering compound to evaluate any significant change in the ELISA result.



Data Summary:

Using a two sided t-test at a 0.05 significance level it was determined that the average Specificity index (SI_{avg}) for the four compounds tested did not differ from 1 (Table 8).

Table 8

Interfering Compound	Conc. (ppm)	SI_{avg}	Significantly different from control by t-test.
Glutamine	100	1.15	No
Glutamic Acid	100	0.89	No
Saxitoxin	20	1.26	No
Kainic Acid	20	1.15	No

Section #5 Linear Range/Limit of Detection/Limit of Quantitation/Sensitivity

Method: Multiple blank mussel tissue samples were spiked with Calbiochem DA standard at the following levels: 2, 3, 5, 10, 15, 20, 30, 40 ug/g then extracted and evaluated by ELISA. To establish the linear range of response the data was evaluated and expressed in Fig. 1. The line of response falls within the bracketed 0.95-1.05 range with data from tissue concentrations from 3 ug/g to 40 ug/g. The data for 2 ug/g falls outside and is not considered within the linear range of the ELISA. The range of assay detection from 3-40 ppm is inclusive of the current NSSP criteria for closing of shellfish beds at 2 mg DA per 100 grams shellfish tissue.

Figure 2 plots the coefficient of variation for each concentration within the linear range which are all under 10%. We can calculate the limit of detection (LOD) of the method and the limit of quantitation (LOQ) shown below using this data.

Data Summary

Linear range of the method as implemented is 3-40 ppm DA in tissue (Fig.1).

The limit of detection (LOD) of the method as implemented is 0.91 ppm DA in tissue.

The limit of quantitation (LOQ) of the method as implemented is 3.0 ppm DA in tissue.

Linear Range Plot – Figure 1

The linear range of the ELISA is established to be from 3 to 40 ppm DA in mussel tissue (Figure 1).



Figure 1: Linearity of Beacon DA ELISA

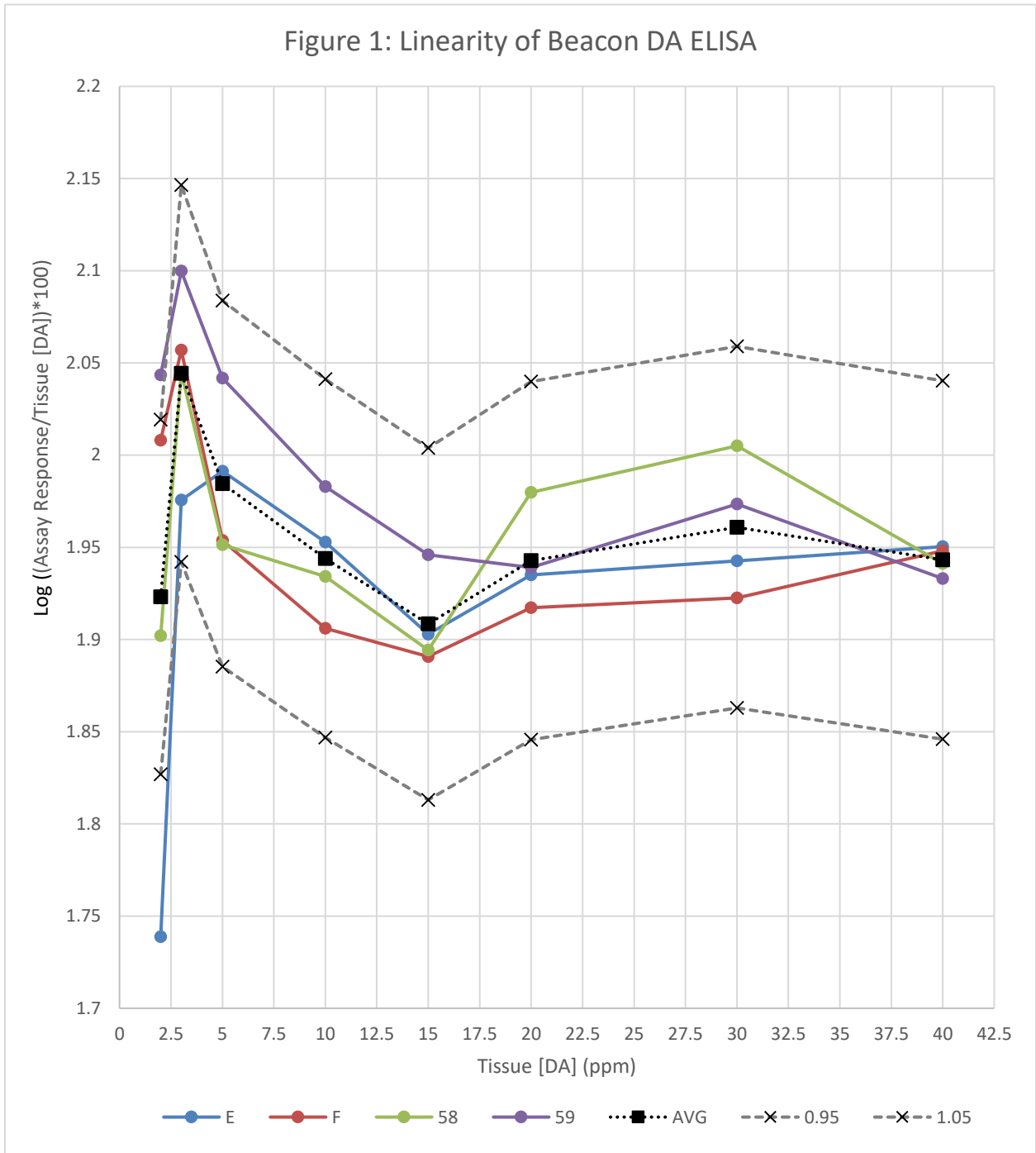
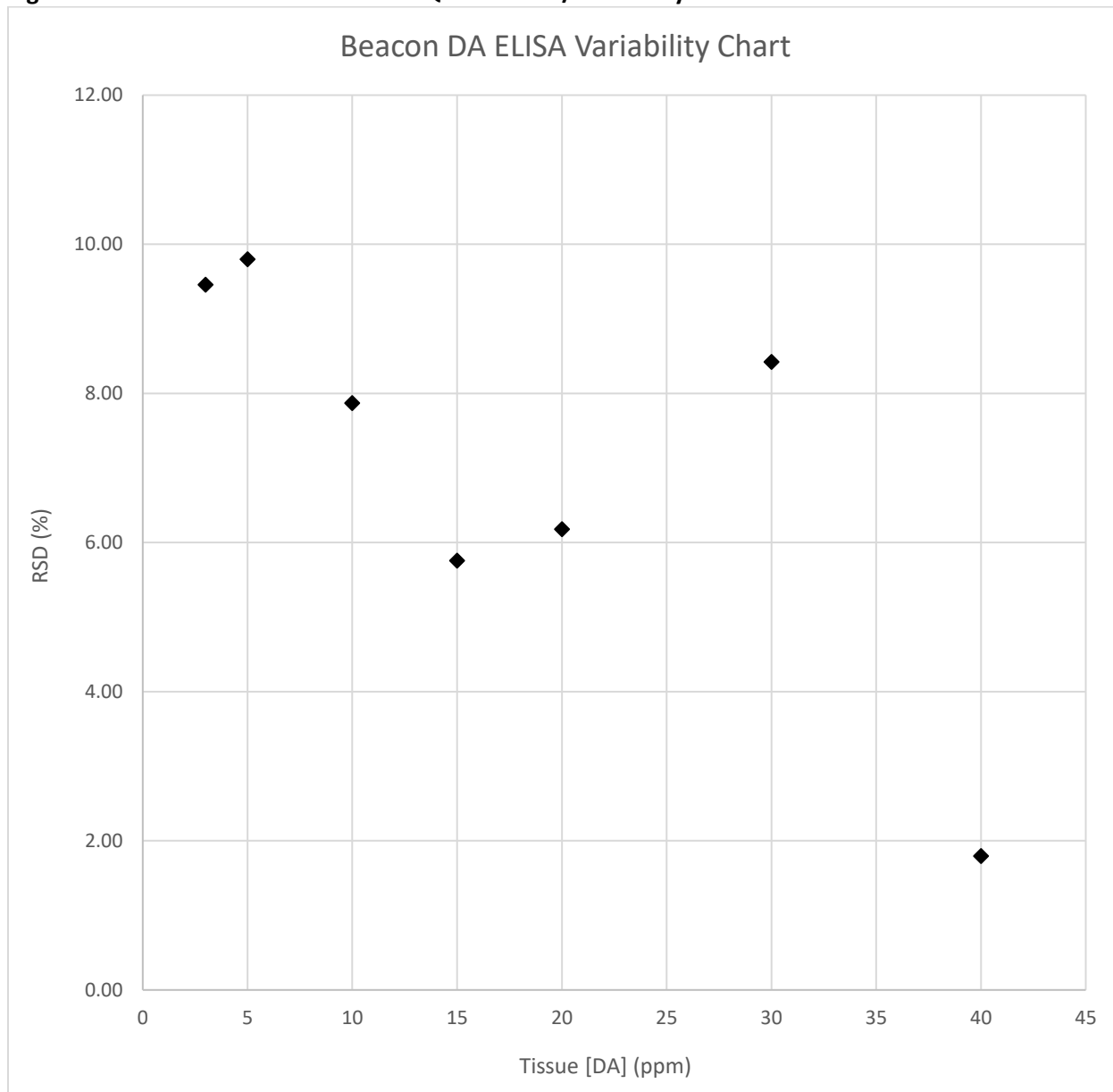




Figure 2: Limit of Detection & Limit of Quantitation/Sensitivity





Section # 6 - Comparability to NSSP Approved Method for Biotoxin Testing –HPLC

Reference from NSSP Guide for the control of Molluscan Shellfish 2015 Revision.

Table 2- Approved Methods for Marine Biotoxin Testing for ASP: M.A. Quilliam, M.Xie and W.R. Hardstaff. 1991. Rapid Extraction and Cleanup Procedure for the Determination of Domoic Acid in Tissue Sample. NRC Institute for Marine Biosciences, Technical Report #64 National Research Council Canada #33001.

Method: The HPLC uses a C-18 reverse phase chromatography column with a mobile phase of 10% acetonitrile and 0.1% trifluoroacetic acid. The mussel tissue extracts have been prepared using the sample extraction procedure for the ELISA which are then diluted 1:5 with 10% acetonitrile prior to injection into the HPLC system. The 2 procedures use the same sample extracts and the results are compared in Table 9. There were 14 mussel tissue samples tested of which 50% were naturally incurred samples and 50% were spiked with Calbiochem DA standard.

Data Summary for the comparison of the new method to the officially recognized method:

Value for the test of symmetry for the data by HPLC reference method: 0.93

Value for the test of symmetry for the data by the DA ELISA method: 1.45

Symmetry is within the range of -2 to +2 and is not a significant degree of skewness.

Variance of data generated by the HPLC reference method: 166.90

Variance of the data generated by the DA ELISA method: 675.73

Ratio of the larger to smaller of the variances: 4.05

This value indicates a lack of homogeneity of variance and indicates the use of a Welch’s t-test for further data analysis to determine if there is a difference between the data means.

Based upon the Welch’s t-test there no significant difference between these two analytical methods.

Table 9

Sample	Collection Date	HPLC Data	DA ELISA
<i>Mytilus edulis</i>		DA (ppm)	DA (ppm)
1	9/6/16	9.48	9.50
2	8/30/16	4.78	4.2
3	8/30/16	16.14	19.80
4	01/30/17	4.42	4.80
5	01/30/17	8.77	8.70
6	01/30/17	15.78	22.80
7	01/30/17	28.49	26.20
8	9/20/16	10.64	21.30
9	10/3/16	27.04	51.40
10	9/20/16	1.60	6.90
11	9/28/16	43.11	91.80
12	9/19/16	17.80	36.70
13	9/26/16	39.79	68.70
14	10/3/16	12.10	22.70



	SKEW	0.93	1.45
	VARIANCE	166.90	675.73
	Ratio of Variance		4.05
	Welch's T-test		-1.43
	df =	(19)	
	T =	2.09	
Conclusion: Means are not different between the 2 methods of analysis.			

Discussion and Summary

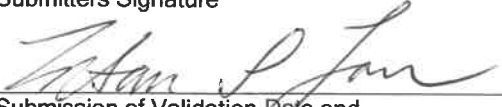
The results of this single laboratory validation demonstrate that the Beacon Domoic Acid (ASP) Plate Kit is an effective procedure for quantitative determination of DA residues in *Mytilus edulis* shellfish tissue. Data presented for ELISA performance meets the validation criteria for accuracy/trueness, measurement uncertainty, ruggedness, precision and recovery. The specificity of the test kit was challenged with four compounds of potential interference and was found to perform properly at DA levels of interest. The linear range of the ELISA was determined to be 3 to 40 ppm which brackets the NSSP established criteria of 20 ppm for the ASP biotoxin in shellfish beds. This linear range would allow for the continued use of the method should a lower criteria be established. The comparative data from the ELISA and the officially recognized HPLC method demonstrate good correlative performance. The ability to use the same sample extract on the ELISA and HPLC confers ease of use for confirmatory testing. The sample throughput is high, while cost and training requirements are minimal. The Beacon Domoic Acid (ASP) Plate Kit is an appropriate tool for quantification of DA residues for use in biotoxin monitoring programs as it allows rapid sample analysis and turnaround time.

ISSC Method Application and Single Lab Validation Checklist For Acceptance of a Method for Use in the NSSP

The purpose of single laboratory validation in the National Shellfish Sanitation Program (NSSP) is to ensure that the analytical method under consideration for adoption by the NSSP is fit for its intended use in the Program. A Checklist has been developed which explores and articulates the need for the method in the NSSP; provides an itemized list of method documentation requirements; and, sets forth the performance characteristics to be tested as part of the overall process of single laboratory validation. For ease in application, the performance characteristics listed under validation criteria on the Checklist have been defined and accompany the Checklist as part of the process of single laboratory validation. Further a generic protocol has been developed that provides the basic framework for integrating the requirements for the single laboratory validation of all analytical methods intended for adoption by the NSSP. Methods submitted to the Interstate Shellfish Sanitation Conference (ISSC) Laboratory Methods Review (LMR) Committee for acceptance will require, at a minimum, six (6) months for review from the date of submission.

Name of the New Method		Domoic Acid (ASP) Plate Kit	
Name of the Method Developer		Dr. Titan Fan	
Developer Contact Information		Beacon Analytical Systems, Inc. 82 Industrial Park Road Saco, ME 04072 Tel. (207) 571-4302 Fax (207) 602 6502 Email: titan@beaconkits.com	
Checklist	Y/N	Submitter Comments	
A. Need for the New Method			
1. Clearly define the need for which the method has been developed.		There is a need for an inexpensive laboratory method with the ability to process multiple samples rapidly and quantify the domoic acid levels in mussel tissue.	
2. What is the intended purpose of the method?		The method can be used in shellfish bed monitoring programs to document the quantifiable levels of domoic acid in mussel tissue.	
3. Is there an acknowledged need for this method in the NSSP?		The method offers higher sample thrupt and quantifiable results to monitor increasing or decreasing levels of domoic acid.	
4. What type of method? i.e. chemical, molecular, culture, etc.		It is an immunochemical method utilizing an antibody specific to the toxin in an enzyme linked immunosorbent assay (ELISA).	
B. Method Documentation			
1. Method documentation includes the following information:			
Method Title		Domoic Acid (ASP) Plate Kit	
Method Scope		Quantitative Analysis of Domoic Acid in mussel tissue	
References		Performance Validation Report and Product Insert included with this submission.	
Principle		Composite mussel samples are extracted and run on an ELISA Test Kit which quantifies Domoic Acid residues by comparison to an internally run calibration curve.	
Any Proprietary Aspects		Yes, this is a commercial test kit.	
Equipment Required		Sample Preparation- Blender, scale, extraction container with lid (10-20 ml), vortex mixer, centrifuge (12,000 rcf), Disposable centrifuge tubes. Calibrated variable pipettes 1.0 ml and 0.010 ml with disposable tips. Plate Reader, timer, wash bottle.	
Reagents Required		Methanol and laboratory grade water in a 1:1 ratio. 10% Acetonitrile in laboratory grade water.	
Sample Collection, Preservation and Storage Requirements		Shellfish should be collected according to standard industry practices and stored at 2-8°C before testing.	
Safety Requirements		Protective safety items are indicated such as safety glasses, gloves and lab coat. Kit reagents including	

		calibrators, conjugates and extracts should be handled with caution since they contain a toxic substance. The kit Stop Solution is a strong acidic solution (1 N Hydrochloric Acid) and needs to be safely handled and disposed of appropriately.
Clear and Easy to Follow Step-by-Step Procedure		Product Insert is included in each test kit and included in this submission package.
Quality Control Steps Specific for this Method		The Certificate of Conformity included with each kit documents the performance characteristics of the Test Kit Lot Reagents. This provides the test operators a reference to evaluate the results generated in their laboratory.
C. Validation Criteria		
1. Accuracy / Trueness		SLV - Section 1
2. Measurement Uncertainty		SLV - Section 1
3. Precision Characteristics (repeatability and reproducibility)		SLV - Section 3
4. Recovery		SLV- Section 3
5. Specificity		SLV- Section 4
6. Working and Linear Ranges		SLV - Section 5
7. Limit of Detection		SLV - Section 5
8. Limit of Quantitation / Sensitivity		SLV - Section 5
9. Ruggedness		SLV- Section 2
10. Matrix Effects		None observed.

11. Comparability (if intended as a substitute for an established method accepted by the NSSP)		SLV - Section 6
D. Other Information		
1. Cost of the Method		The price per sample is eight to nine dollars dependent upon the number of samples tested during one ELISA run, and/or the volume of kits purchased.
2. Special Technical Skills Required to Perform the Method		Knowledge of GMP laboratory skills as well as proper pipetting technique, and safe handling of solvents.
3. Special Equipment Required and Associated Cost		An ELISA Plate Reader is required which can range in price from a low cost unit at approximately \$2,600 to a higher cost of \$15,000 USD unit depending upon complexity.
4. Abbreviations and Acronyms Defined		ASP-amnesic shellfish poisoning. DA-domoic acid. ELISA-Enzyme Linked Immunosorbent Assay. OD-Optical Density (Reader Output). HRP-horse radish peroxidase enzyme. % B/B0- percent of measured bound fraction in a test sample divided by the total bound from a blank Calibrator. ppm = parts per million, equivalent to mg/kg, ppb –parts per billion.
5. Details of Turn Around Times (time involved to complete the method)		One assay can be completed in under 90 minutes including sample preparation (12 samples). One plate can be used to test a maximum of 36 samples.
6. Provide Brief Overview of the Quality Systems Used in the Lab		<p>Beacon Analytical Systems is ISO 9001:2015 approved for their quality systems for Immunochemical test kit development, manufacturing and supporting activities.</p> <p><u>Overview of Kit Quality Control</u> - Each kit is tested following the ELISA procedure in the product insert. During manufacturing operation duplicates of the Kit Calibrator Reagents are run in order to meet established criteria prior to shipment.</p> <p>Each Domoic Acid (DA) Calibrator's absorbance (OD) and binding characteristics (% B/B0) must be within a set of ranges. Ranges have been established for the Domoic Acid Plate Kit from historical data.</p> <p>All plate components are tested for precision prior to using them in kits. The tolerance for variation within one lot of plates is less than or equal to 5%.</p> <p>-DA Calibration solutions are prepared using certified reference standard material purchased from the Canadian National Resource Council and are tested to be within 2% of the previous lot of control.</p> <p>- The R² correlation of the DA Kit Calibration Curve should be 0.99 or above.</p> <p>- All CV's must be less than or equal to 5%.</p> <p>- All QC data is kept electronically and backed up with hard copies at our manufacturing plant.</p>
Submitters Signature		Date: 06/30/2017
		Date:
Submission of Validation Data and Draft Method to Committee		Date:
Reviewing Members		Date:

Accepted	Date:
Recommendations for Further Work	Date:
Comments:	

DEFINITIONS

1. **Accuracy/Trueness** - Closeness of agreement between a test result and the accepted reference value.
2. **Analyte/measurand** - The specific organism or chemical substance sought or determined in a sample.
3. **Blank** - Sample material containing no detectable level of the analyte or measurand of interest that is subjected to the analytical process and monitors contamination during analysis.
4. **Comparability** - The acceptability of a new or modified method as a substitute for an established method in the NSSP. Comparability must be demonstrated for each substrate or tissue type by season and geographic area if applicable.
5. **Fit for purpose** - The analytical method is appropriate to the purpose for which the results are likely to be used.
6. **HORRAT value** - HORRAT values give a measure of the acceptability of the precision characteristics of a method.⁴
7. **Limit of Detection** - the minimum concentration at which the analyte or measurand can be identified. Limit of detection is matrix and analyte/measurand dependent.⁴
8. **Limit of Quantitation/Sensitivity** - the minimum concentration of the analyte or measurand that can be quantified with an acceptable level of precision and accuracy under the conditions of the test.
9. **Linear Range** - the range within the working range where the results are proportional to the concentration of the analyte or measurand present in the sample.
10. **Measurement Uncertainty** - A single parameter (usually a standard deviation or confidence interval) expressing the possible range of values around the measured result within which the true value is expected to be with a stated degree of probability. It takes into account all recognized effects operating on the result including: overall precision of the complete method, the method and laboratory bias and matrix effects.
11. **Matrix** - The component or substrate of a test sample.
12. **Method Validation** - The process of verifying that a method is fit for purpose.¹
13. **Precision** - the closeness of agreement between independent test results obtained under stipulated conditions.^{1, 2} There are two components of precision:
 - a. **Repeatability** - the measure of agreement of replicate tests carried out on the same sample in the same laboratory by the same analyst within short intervals of time.
 - b. **Reproducibility** - the measure of agreement between tests carried out in different laboratories. In single laboratory validation studies reproducibility is the closeness of agreement between results obtained with the same method on replicate analytical portions with different analysts or with the same analyst on different days.

14. **Quality System** - The laboratory's quality system is the process by which the laboratory conducts its activities so as to provide data of known and documented quality with which to demonstrate regulatory compliance and for other decision-making purposes. This system includes a process by which appropriate analytical methods are selected, their capability is evaluated, and their performance is documented. The quality system shall be documented in the laboratory's quality manual.
15. **Recovery** - The fraction or percentage of an analyte or measurand recovered following sample analysis.
16. **Ruggedness** - the ability of a particular method to withstand relatively minor changes in analytical technique, reagents, or environmental factors likely to arise in different test environments.⁴
17. **Specificity** - the ability of a method to measure only what it is intended to measure.¹
18. **Working Range** - the range of analyte or measurand concentration over which the method is applied.

REFERENCES:

1. Eurachem Guide, 1998. The Fitness for Purpose of Analytical Methods. A Laboratory Guide to Method Validation and Related Topics. LGC Ltd. Teddington, Middlesex, United Kingdom.
2. IUPAC Technical Report, 2002. Harmonized Guidelines for Single-Laboratory Validation of Methods of Analysis, Pure Appl. Chem., Vol. 74, (5): 835-855.
3. Joint FAO/IAEA Expert Consultation, 1999. Guidelines for Single-Laboratory Validation of Analytical Methods for Trace-Level Concentrations of Organic Chemicals.
4. MAF Food Assurance Authority, 2002. A Guide for the Validation and Approval of New Marine Biotxin Test Methods. Wellington, New Zealand.
5. National Environmental Laboratory Accreditation. , 2003. Standards. June 5.
6. EPA. 2004. EPA Microbiological Alternate Procedure Test Procedure (ATP) Protocol for Drinking Water, Ambient Water, and Wastewater Monitoring Methods: Guidance. U.S. Environmental Protection Agency (EPA), Office of Water Engineering and Analysis Division, 1200 Pennsylvania Avenue, NW, (4303T), Washington, DC 20460. April.



Domoic Acid (ASP)

Plate Kit

Cat. # 20-0249

Product Insert

PLEASE READ COMPLETELY BEFORE USE

INTENDED USE

The Beacon Domoic Acid (ASP) Plate Kit is a competitive ELISA for the quantitative analysis of domoic acid in shellfish samples.

USE PRINCIPLES

The Beacon Domoic Acid (ASP) plate kit is a competitive enzyme-labeled immunoassay for the biotoxin which causes Amnesic Shellfish Poisoning (ASP). Shellfish sample extract(s) or calibrator solution(s) are pipetted into a test well followed by Domoic Acid HRP enzyme conjugate to initiate the reaction. During a 30 minute incubation period, domoic acid from the sample and domoic acid HRP enzyme conjugate compete for binding to the domoic acid antibody coated on the plate wells. Following this incubation, the wells are washed to remove any unbound domoic acid and HRP enzyme conjugate. After washing, a colorless substrate is added to the wells and any bound enzyme conjugate will convert the substrate to a blue color. Following another 30 minute incubation, the reaction is stopped with the addition of stop solution and the amount of color in each well is measured. The color of the unknown sample is compared to the color of the calibrators and the domoic acid concentration of the sample is derived. The color intensity is inversely proportional to the amount of domoic acid present.

MATERIALS PROVIDED

The kit in its original packaging can be used until the end of the month indicated on the box label when stored at 2 to 8 °C.

- **Plate** – (1) containing 12 test strips of 8 wells each vacuum-packed in aluminized pouch with indicating desiccant.
- **Domoic Acid Calibrators**– (4) vials each containing 2 ml with a concentration of 0, 0.5, 5, and 50 µg/L (ppb) Domoic Acid
- **Domoic Acid HRP Enzyme Conjugate** – (1) vial containing 12 ml
- **Substrate** – (1) vial containing 14 ml
- **Stop Solution** – (1) vial containing 14 ml (Caution! Contains 1N HCl. Handle with care.)
- **Product Insert** containing instructions for use.
- **Certificate of Conformity** (Specific to each Kit Lot#).

MATERIALS REQUIRED BUT NOT PROVIDED

Acetonitrile, & Methanol (ACS grade)	Timer
Laboratory quality distilled or deionized water	Wash bottle
Variable volume pipettes with disposable tips capable of dispensing 10-100 microliters (µl), and up to 1000 µl.	Vortex mixer
Multi-channel pipette; 8 channel capable of dispensing 100 µl	Paper towels or equivalent absorbent material
Microwell plate or strip reader with 450 nm filter	Disposable micro centrifuge tubes
Microcentrifuge capable of a speed of 12,000 rcf. (x g)	Kitchen Blender for sample homogenization

SPECIFICITY

Domoic Acid (DA) is an amino acid similar in structure to kainic acid which naturally occurs in some seaweed. The % cross reactivity of several compounds relative to DA is shown in the table below.

Compound	% CR	Compound	% CR
Domoic acid	100 %	Saxitoxin	< 0.1 %
Glutamine	< 0.1 %	Kainic acid	0.005 %
Glutamic acid	< 0.1 %		

KIT HANDLING NOTES and PRECAUTIONS

- Store all kit components at 4 °C to 8 °C (39 °F to 46 °F) when not in use.
- Each reagent is optimized for use in the Beacon Domoic Acid (ASP) Plate Kit. Do not substitute reagents from any other manufacturer into the test kit. Do not combine reagents from other Beacon Plate Kits with different lot numbers.
- Dilution or adulteration of reagents or samples not called for in the procedure may result in inaccurate results.
- Do not use reagents after expiration date.
- Reagents should be brought to room temperature (RT), 20 to 28 °C (62 to 82 °F) prior to use. Avoid prolonged (> 24 hours) storage at room temperature.
- Domoic acid calibrators contain 10% acetonitrile and should be kept tightly capped to minimize evaporation.
- The Stop Solution is 1N hydrochloric acid, which is corrosive and an irritant. Avoid contact with skin and mucous membranes. Immediately clean up any spills and wash area with copious amounts of water. If contact should occur, immediately flush with copious amounts of water.
- Precise transfer of samples and reagents by using an appropriate and calibrated pipette is critical to obtain proper assay results. Please pipette carefully.
- If running more than two strips at once, the use of a multichannel pipette is required.
- In one assay a maximum of four strips (32 wells) is recommended, for example –4 calibrators in duplicate (8 wells), and 12 sample extracts in duplicate (24 wells).

SAMPLE DILUTION BUFFER PREPARATION- 10% ACETONITRILE/Water

- Mix 1 part ACS grade acetonitrile with 9 parts distilled or deionized (DI) water to make the Sample Dilution Buffer for the shellfish samples.
- Transfer to a clean glass container with tight-fitting lid and swirl to mix. Store tightly sealed to minimize evaporation.

SAMPLE EXTRACTION BUFFER PREPARATION- 50% METHANOL/Water

- Mix 1 part ACS grade methanol with 1 part distilled or deionized (DI) water to make the Sample Extration Buffer.
- Transfer to a clean glass container with tight-fitting lid and swirl to mix. Store tightly sealed to minimize evaporation.

SAMPLE PREPARATION - Shellfish Tissue Extract –*Mytilus edulis* (Blue Mussel)

1. Remove shellfish tissue (12-15 animals) from shell, wash, drain dry and homogenize using a kitchen blender.
2. Weigh 2 g of homogenized tissue and add 8 ml of a 50% Methanol/Water solution.
3. Mix for 3 minutes using Vortex mixer (4 X dilution)
4. Transfer 1 ml into a microcentrifuge tube and centrifuge at 12,000 x g for 5 minutes. Extracts can be stored at -20°C.
5. Prepare a 1:1000 dilution of the supernatant with Sample Dilution Buffer using the following procedure:
A. 1:10 dilution - 50 microliters of supernatant layer avoiding any particulates, into 450 microliters Sample Dilution Buffer, Mix.
B. 1:100 dilution – 10 microliters of dilution **A.** into 990 microliters Sample Dilution Buffer, Mix,
6. Use **B.** in ELISA. – Total Dilution Factor (TDF) = 4000

Shellfish Analysis:

- ✓ EU Screening Level = 20 ppm (20 mg/kg) Assay Dilution Factors are set to detect 20 ppm Domoic Acid,

Extraction of Shellfish Tissue and Preparation for ELISA	
Dilution of shellfish homogenate in water (2 g homogenized tissue with 8 ml 50% MEOH/DI H ₂ O)	4 X Dilution
*Secondary Dilution into Sample Dilution Buffer	1000 X Dilution
<i>Total Dilution Factor</i> (TDF) to obtain Tissue Levels of Domoic Acid	4000 X
✓ Assay Range of Detection in Tissue	2 mg / kg to 200 mg / kg
Domoic Acid Plate Kit Calibrators ug / L (ppb)	Predicted Tissue Levels (X 4000 TDF)
Negative Control (Blank)	0
0.5	2 ppm
5.0	20 ppm
50.0	200 ppm

ASSAY PROCEDURE

(Note: Running Calibrators and samples in duplicate will provide optimal assay precision and accuracy.)

1. Allow reagents and sample extracts to reach RT prior to running the test.
2. Place the appropriate number of test wells into a micro well holder. Be sure to re-seal unused wells in the zip-lock bag with desiccant.
3. Using a pipette with disposable tips, dispense **100 µl** of the **Calibrator or sample extract** into the appropriate test wells. Please use a clean pipette tip for each sample addition.
4. Dispense **100 µl** of the **HRP Enzyme Conjugate** into each well.
5. Shake the plate gently for 30 seconds using a back and forth motion. Then incubate the wells for **30 minutes** at RT.
6. Decant the contents of the wells into an appropriate waste container. Fill the wells to overflowing with laboratory grade water and then decant. Repeat four times for a total of five washes.
7. Following the last wash, tap the inverted wells onto absorbent paper to remove the last of the water.
8. Dispense **100 µl** of the **Substrate** into each well. Shake the plate gently for 30 seconds using a back and forth motion.
9. Incubate the wells for **30 minutes** at RT.
10. Dispense **100 µl of the Stop Solution** into each well.
11. Measure and record the absorbance (Optical Density; OD) of the wells at 450 nm using a strip or plate reader.
12. To obtain the concentration of Domoic acid in the sample multiply the results by the Total Dilution Factor of 4000.

Note: If the sample absorbance is higher or lower than the 0.5 or 50 ppb Calibrator results, the tissue levels should be expressed as less than or greater than the corresponding tissue levels (<2ppm or >200ppm DA). The sample dilution can be modified appropriately and retested along with another set of Calibrators.

CALCULATE RESULTS

1. Semi-quantitative results can be derived visually by simple comparison of the sample color to the color of the Calibrator wells. Samples containing less color than a Calibrator will have a concentration of Domoic Acid greater than the tissue correlated concentration of the Calibrator. Samples containing more color than a Calibrator will have a concentration less than the tissue correlated concentration of the Calibrator.
2. It is preferred for quantitative results to be determined using commercially available software for ELISA evaluation such using a 4-Parameter curve fit. Alternatively, a semi-log curve fit can be used if 4-Parameter software is not available. Samples with OD's greater than the lowest calibrator, or lower than the highest calibrator will need to be diluted accordingly and repeated with and calibrators in an additional run.
3. Beacon can supply a spreadsheet template which can be used for data reduction. Please contact Beacon for further details.

SAMPLE CALCULATIONS

Well Contents	OD	Average OD ± SD*	%RSD	%B/Bo**
Negative Control	2.033 1.994	2.014 ± 0.027	1.4	100
0.5 ppb Calibrator	1.610 1.671	1.640 ± 0.043	2.7	81
5 ppb Calibrator	1.095 1.155	1.125 ± 0.042	3.8	56
50 ppb Calibrator	0.501 0.482	0.492 ± 0.013	2.7	24

Actual values may vary; this data is for example purposes only.

* Standard deviation

**B/Bo% equals the average sample absorbance divided by the average 0 ppb Calibrator absorbance multiplied by 100.

TECHNICAL ASSISTANCE

For questions regarding this kit or for additional information about Beacon products, call (207) 571-4302 or contact us at info@beaconkits.com.

Safety- To receive complete safety information on this product, contact Beacon Analytical Systems, Inc. and request Safety Data Sheets. Stop Solution is 1N hydrochloric acid. Handle with care.

General Limited Warranty

Beacon Analytical Systems, Inc. ("Beacon") warrants the products manufactured by it against defects in materials and workmanship when used in accordance with the applicable instructions for a period not to extend beyond a product's printed expiration date. BEACON MAKES NO OTHER WARRANTY, EXPRESSED OR IMPLIED. THERE IS NO WARRANTY OF MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE. The warranty provided herein and the data, specifications and descriptions of Beacon products appearing in published catalogues and product literature may not be altered except by express written agreement signed by an officer of Beacon. Representations, oral or written, which are inconsistent with this warranty or such publications are not authorized and, if given, should not be relied upon.

In the event of a breach of the foregoing warranty, Beacon's sole obligation shall be to repair or replace, at its option, any product or part thereof that proves defective in materials or workmanship within the warranty period, provided the customer notifies Beacon promptly of any such defect. The exclusive remedy provided herein shall not be deemed to have failed of its essential purpose so long as Beacon is willing and able to repair or replace any nonconforming Beacon product or part. Beacon shall not be liable for consequential, incidental, special or any other indirect damages resulting from economic loss or property damage sustained by a customer from the use of its products. However, in some states the purchaser may have rights under state law in addition to those provided by this warranty.

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PUBLIC HEALTH SERVICE U.S. FOOD AND DRUG ADMINISTRATION OFFICE OF FOOD SAFETY SHELLFISH AND AQUACULTURE POLICY BRANCH 5001 CAMPUS DRIVE COLLEGE PARK, MD 20740-3835 TEL. 240-402-2151/2055/4960 FAX 301-436-2601 CFSANDSSLEOS@FDA.HHS.GOV		
SHELLFISH LABORATORY EVALUATION CHECKLIST		
LABORATORY:		
ADDRESS:		
TELEPHONE:	FAX:	EMAIL:
DATE OF EVALUATION:	DATE OF REPORT:	LAST EVALUATION:
LABORATORY REPRESENTED BY:		TITLE:
LABORATORY EVALUATION OFFICER:		SHELLFISH SPECIALIST:
OTHER OFFICIALS PRESENT:		TITLE:
Items which do not conform are noted by:		Conformity is noted by a “√”
C- Critical K - Key O - Other NA- Not Applicable		
Check the applicable analytical methods:		
<input type="checkbox"/>	Preparation of Samples for the Alkaline Phosphatase Probe Method: Direct Plating [PART III]	
<input type="checkbox"/>	Preparation of Samples for the Alkaline Phosphatase Probe Method: APW Enrichment and Colony Isolation [PART III]	
<input type="checkbox"/>	Alkaline Phosphatase Probe Hybridization [PART III] - Note: Temperature ranges for hybridization can be expanded with an appropriate study as described in the method.	

PART I – QUALITY ASSURANCE			
ITEM			
Code	REF		
1.1 Quality Assurance (QA) Plan			
K	4, 6	<input type="checkbox"/>	1.1.1 Written Plan (check those items which apply).
		<input type="checkbox"/>	a. Organization of the laboratory.
		<input type="checkbox"/>	b. Staff training requirements.
		<input type="checkbox"/>	c. Standard operating procedures.
		<input type="checkbox"/>	d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance and rejection criteria established.
		<input type="checkbox"/>	e. Laboratory safety.
		<input type="checkbox"/>	f. Internal performance assessment.
		<input type="checkbox"/>	g. External performance assessment.
C	4	<input type="checkbox"/>	1.1.2 The QA plan is implemented.
K	6	<input type="checkbox"/>	1.1.3 The Laboratory participates in a Vibrio proficiency testing program annually. Specify the program(s): _____
1.2 Educational/Experience Requirements			
C	State's Human Resources Department	<input type="checkbox"/>	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	<input type="checkbox"/>	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology or equivalent discipline with at least two (2) years of laboratory experience.
K	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three (3) months of experience in laboratory sciences.
1.3 Work Area			
O	4, 6	<input type="checkbox"/>	1.3.1 Adequate for workload and storage.
K	6	<input type="checkbox"/>	1.3.2 Clean, well-lighted.
K	6	<input type="checkbox"/>	1.3.3 Adequate temperature control.
O	6	<input type="checkbox"/>	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	6	<input type="checkbox"/>	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained
1.4 Laboratory Equipment			
K	5	<input type="checkbox"/>	1.4.1 To determine the pH of prepared media and reagents, the pH meter has a standard accuracy of at least 0.1 pH units.

K	9	<input type="checkbox"/>	1.4.2 The pH electrodes being used consist of a pH half-cell and reference half-cell or equivalent combination electrode free from Ag/AgCl or contains an ion exchange barrier preventing passage of Ag ions into the solution which may affect the accuracy of the pH reading.
K	6	<input type="checkbox"/>	1.4.3 The effect of temperature on the pH is compensated for by an internal/external ATC probe or by manual adjustment.
K	4	<input type="checkbox"/>	1.4.4 The pH meter is calibrated daily or with each use. Results are recorded and records maintained.
K	6	<input type="checkbox"/>	1.4.5 A minimum of two (2) standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second is near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
K	4, 17	<input type="checkbox"/>	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt procedure or through determination of the slope (<i>Circle the method used</i>).
K	5, 15	<input type="checkbox"/>	1.4.7 The balances used provide a sensitivity of at least 0.01 g at the weights of use for direct plating and 0.1 g for MPN.
K	6	<input type="checkbox"/>	1.4.8 Balance calibrations are checked monthly according to manufacturer specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance calibrations is verified at the weight range of use. Results are recorded and records maintained.
K	6	<input type="checkbox"/>	1.4.9 Refrigerator temperatures are monitored at least once daily on workdays. Results are recorded and records maintained.
C	12, 15	<input type="checkbox"/>	1.4.10 Refrigerator temperatures in which AP-probes are stored are maintained between 2 and 8 °C.
K	1	<input type="checkbox"/>	1.4.11 The temperature of general purpose refrigerators, those not containing AP-probes, are maintained between 0 and 4 °C.
C	2	<input type="checkbox"/>	1.4.12 Freezer temperatures are maintained at -15 °C or below.
K	6	<input type="checkbox"/>	1.4.13 Freezer temperature is monitored at least once daily on workdays. Results are recorded and records maintained.
C	12	<input type="checkbox"/>	1.4.14 The temperature of the incubator is maintained at 35 ± 2.0 °C.
C	6	<input type="checkbox"/>	1.4.15 Working thermometers used in the air incubators are graduated in at least 0.5 °C increments.
K	5, 8	<input type="checkbox"/>	1.4.16 Working thermometers are located on top and bottom shelves of use in the air incubator or appropriately placed based on the results of spatial temperature checks.
C	6	<input type="checkbox"/>	1.4.17 Temperature of the water bath is maintained appropriately under all loading conditions.
C	5	<input type="checkbox"/>	1.4.18 Working thermometers used in the water bath are graduated in at least 0.1 °C increments.
K	4, 6	<input type="checkbox"/>	1.4.19 Air incubator/water bath temperatures are taken twice daily on workdays. Results are recorded and records maintained.
C	3	<input type="checkbox"/>	1.4.20 All working thermometers are appropriately immersed.

C	5	<input type="checkbox"/>	1.4.21 Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	5, 6	<input type="checkbox"/>	1.4.22 A standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35, 42, 54 and/or 55 °C (54 °C for <i>Vibrio parahaemolyticus</i> and 55 °C for <i>Vibrio vulnificus</i>). These calibration records (certificates of calibration) are maintained.
K	3	<input type="checkbox"/>	1.4.23 Standards thermometers are checked annually for accuracy by ice point determination. Results are recorded and maintained. Date of most recent determination: .
C	5	<input type="checkbox"/>	1.4.24 Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with at least an accuracy of ±0.05 °C are used as the laboratory standards thermometer (<i>Circle the thermometer type used</i>).
K	3, 8	<input type="checkbox"/>	1.4.25 All working thermometers are checked annually against the standards thermometer at the temperature(s) of use. Results for are recorded and records maintained. <u>The accuracy of working thermometers is checked annually against the standards thermometer either at the temperatures at which they are used or by ice point determination. Results are recorded and records maintained.</u>
O	8	<input type="checkbox"/>	1.4.26 Appropriate pipet aids are available and used to inoculate samples.
K	7	<input type="checkbox"/>	1.4.27 Micropipettors are calibrated annually and checked for accuracy quarterly at volumes of use. Results are recorded and records maintained.
			1.5 Labware and Glassware Washing
K	5	<input type="checkbox"/>	1.5.1 Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding material.
K	5	<input type="checkbox"/>	1.5.2 Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and sample.
O	5	<input type="checkbox"/>	1.5.3 Dilution bottles and tubes are made of borosilicate glass or plastic and closed with secure caps or screw caps with nontoxic liners.
K	5	<input type="checkbox"/>	1.5.4 Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method of preparation is used to ensure the appropriate volumes of diluent.
C	5	<input type="checkbox"/>	1.5.5 Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1 mL aliquots; nor, are pipettes larger than 1.1 mL used to deliver 0.1 mL aliquots.

K	5	<input type="checkbox"/>	1.5.6 In washing reusable pipets, glassware and labware, a succession of at least three (3) fresh water rinses plus a final rinse of deionized water is used to thoroughly rinse off all detergent.
C	8	<input type="checkbox"/>	1.5.7 An alkaline or acidic detergent is used for washing glassware/labware.
C	6	<input type="checkbox"/>	1.5.8 With each load of labware/glassware washed, the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue (BTB) solution. Results are recorded, and records maintained.
			1.6 Sterilization and Decontamination
K	5	<input type="checkbox"/>	1.6.1 The autoclave is of sufficient size to accommodate the workload.
K	4	<input type="checkbox"/>	1.6.2 Routine autoclave maintenance is performed, and the records are maintained.
C	<u>19, 20, 21</u> <u>6, 8</u>	<input type="checkbox"/>	1.6.3 <u>The autoclave provides sterilization conditions suitable to the load contents. Sterilization temperature range may be 119°C - 124°C as determined by the lab's equipment Quality Assurance Verification Testing and recommended practices from the media manufacturer. Sterilization is determined for each load using a working-maximum registering thermometer, or an appropriate working temperature monitoring device.</u>
K	2, 5, 6	<input type="checkbox"/>	1.6.4 An autoclave standards thermometer (or data logger) has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121 °C. If in-house checks for accuracy of the standards thermometer will be conducted at the steam point, calibration of the autoclave standards thermometer at 100 °C is also recommended, but not required.
K	2, 10, 18	<input type="checkbox"/>	1.6.5 The autoclave standards thermometer (or data logger) is checked every five (5) years for accuracy at either 121 °C by a qualified calibration laboratory; or, is checked in-house at the steam point (100 °C) if it has been previously calibrated at both 100 °C and 121 °C. Any change in temperature at the steam point changes the calibrated temperature at 121 °C by the same magnitude. Date of most recent determination: _____
K	2, 8	<input type="checkbox"/>	1.6.6 Working autoclave thermometers (or data loggers) are checked against the autoclave standards thermometer at 121 °C yearly. Date of last check: _____ Method: _____
K	6	<input type="checkbox"/>	1.6.7 Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded, and the records maintained.
O	6	<input type="checkbox"/>	1.6.8 Heat sensitive tape is used with each autoclave batch.

K	6, 8	<input type="checkbox"/>	<p>1.6.9 Autoclave sterilization records including the length of sterilization cycle, total heat exposure time and chamber temperature are maintained.</p> <p>Type of record: Autoclave log, computer printout or chart recorder tracings. (<i>Circle the appropriate type or types</i>)</p>
K	5, 8	<input type="checkbox"/>	<p>1.6.10 For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180 °C.</p>

K	8	<input type="checkbox"/>	1.6.11 Records of temperature and exposure times are maintained for the operation of the hot-air sterilizing oven during use.
K	8	<input type="checkbox"/>	1.6.12 Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Results are recorded, and records maintained.
K	5	<input type="checkbox"/>	1.6.13 Reusable pipets are stored and sterilized in aluminum or stainless-steel containers.
K	5	<input type="checkbox"/>	1.6.14 Reusable pipets (in canisters) are sterilized in a hot-air oven at 170 °C for two (2) hours.
C	2	<input type="checkbox"/>	1.6.15 The sterility of reusable pipets is determined with each load sterilized. Results are recorded, and records maintained.
C	2	<input type="checkbox"/>	1.6.16 The sterility of autoclave sterilized disposable pipet tips and microcentrifuge tubes is determined with each load sterilized. Results are recorded, and records maintained.
C	2	<input type="checkbox"/>	1.6.17 The sterility of pre-sterilized disposable pipettes, pipet tips and microcentrifuge tubes is determined with each lot received. Results are recorded, and records maintained.
K	8	<input type="checkbox"/>	1.6.18 Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
			1.7 Media and Reagent Preparation
C	12, 15	<input type="checkbox"/>	1.7.1 Media and reagents are prepared from the individual components and pH adjusted appropriately, except in the case of TCBS, which is commercially dehydrated.
K	1, 5, 8	<input type="checkbox"/>	1.7.2 Dehydrated media, and media and reagent components are properly stored in a cool, clean, dry place.
K	1	<input type="checkbox"/>	1.7.3 Media and components are labeled with the analyst's initials, date of receipt, date opened or date of preparation, if applicable (dye solutions).
C	1, 2, 7	<input type="checkbox"/>	1.7.4 Caked or expired media or components are discarded.
C	6	<input type="checkbox"/>	1.7.5 Reagent water is distilled or deionized (<i>circle appropriate choice</i>), tested monthly and exceeds 0.5 megohms-cm resistivity (2 megohms-cm in-line) or is less than 2.0 μSiemens/cm conductivity at 25 °C. (<i>Circle the appropriate water quality descriptor determined</i>). Results are recorded and the records maintained.
C	6	<input type="checkbox"/>	1.7.6 Reagent water for media and diluent preparation is analyzed for residual chlorine monthly and is at a non-detectable level (≤ 0.1 mg/L). Results are recorded, and records maintained. Specify method of determination:
K	6	<input type="checkbox"/>	1.7.7 Reagent water for media and diluent preparation contains <100 CFU/mL as determined monthly using the heterotropic plate count method. Results are recorded, and records maintained.
K	12	<input type="checkbox"/>	1.7.8 The volume and concentration of media (APW) in the tube is suitable for the amount of sample inoculated.
C	2	<input type="checkbox"/>	1.7.9 The total time of exposure of the sugar containing agar VVA to autoclave temperatures does not exceed 45 minutes. Total exposure time of APW and T1N3 agar does not exceed 60 minutes. TCBS, CC and mCPC are not autoclaved.

C	1	<input type="checkbox"/>	1.7.10 Media and diluent sterility is determined for each load sterilized. Results are recorded, and records maintained.
C	1	<input type="checkbox"/>	<p>1.7.11 Media productivity is determined using media-appropriate positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is made from its individual components.</p> <p>Positive <i>Vibrio parahaemolyticus</i> productivity control _____</p> <p>Negative <i>Vibrio parahaemolyticus</i> productivity control _____</p> <p>Positive <i>Vibrio vulnificus</i> productivity control _____</p> <p>Negative <i>Vibrio vulnificus</i> productivity control _____</p>
C	6, 12	<input type="checkbox"/>	1.7.12 The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer requirements and/or method tolerance. Results are recorded, and records are maintained.
			1.8 Storage of Prepared Culture Media and Reagents
K	5	<input type="checkbox"/>	1.8.1 Prepared culture media are stored in a cool, clean, dry place where excessive evaporation and the danger of contamination is minimized.
K	2	<input type="checkbox"/>	1.8.2 Stored media are labeled with the storage expiration date or sterilization date.
K	2	<input type="checkbox"/>	1.8.3 Storage of prepared culture media at room temperature does not exceed seven (7) days.
K	6	<input type="checkbox"/>	1.8.4 Storage under refrigeration of prepared agar plates in sealed plastic bags shall not exceed two (2) weeks.
K	6	<input type="checkbox"/>	1.8.5 Storage under refrigeration of prepared broth media with loose fitting closures shall not exceed one (1) month.
K	6	<input type="checkbox"/>	1.8.6 Storage under refrigeration of prepared broth media and diluent with screw-cap closures shall not exceed three (3) months.
K	12, 15	<input type="checkbox"/>	1.8.7 Refrigerated prepared plates are dried inverted before use to permit the sample to be completely absorbed into the medium to prevent colony spreading, for direct plating.
K	2, 6	<input type="checkbox"/>	1.8.8 All prepared broth media and diluent stored under refrigeration are warmed to room temperature prior to use, at temperatures that do not exceed the medium's incubation temperature.
K	15	<input type="checkbox"/>	1.8.9 Storage at room temperature of Lysis Solution, Ammonium Acetate Buffer, 20XSSC, 1XSSC/SDS, and 3XSSC/SDS for the hybridization procedure shall not exceed three (3) months.
K	15	<input type="checkbox"/>	1.8.10 Storage under refrigeration of Hybridization Buffer for the hybridization procedure shall not exceed one (1) week.

C	15	<input type="checkbox"/>	1.8.11 NBT/BCIP solution and 1XSSC for the hybridization procedure should be made fresh the day of use.
PART II – SHELLFISH SAMPLES			
			2.1 Sample Handling and Receipt
C	1, 5, 12, 15	<input type="checkbox"/>	2.1.1 A representative sample is collected and a chain of custody documenting the history of the sample(s) from collection to final disposal has been established.
K	5, 15	<input type="checkbox"/>	2.1.2 Shellfish samples are received in clean, waterproof, puncture resistant containers loosely sealed or are rejected for regulatory analysis.
K	1, 5	<input type="checkbox"/>	2.1.3 Samples are received labeled with the collector's (or if PHP, company/processor and collector's) name, the source, the time and date of collection or are rejected for regulatory analysis.
C	5, 12, 15	<input type="checkbox"/>	2.1.4 Immediately after collection, samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10 °C with ice or cold packs for transport to the laboratory or rejected. Direct contact of the shellfish with ice in the transport container should be avoided. Once received, the samples are placed under refrigeration unless processed immediately.
K	5, 15	<input type="checkbox"/>	2.1.5 If ice is used in sample transport, samples are rejected if melt water has come in contact with the samples.
C	15	<input type="checkbox"/>	2.1.6 Analysis of the samples is initiated as soon as possible after collection, but not to exceed 36 hours. If processing IQF samples, samples are defrosted under refrigeration for no longer than 36 hours once removed from the freezer.
			2.2 Preparation of Samples for Analysis
K	2, 11	<input type="checkbox"/>	2.2.1 Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2, 11	<input type="checkbox"/>	2.2.2 Blades of shucking knives are not corroded.
K	5, 11	<input type="checkbox"/>	2.2.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2, 11	<input type="checkbox"/>	2.2.4 The faucet used for rinsing the shellfish does not contain an aerator.
K	5, 11	<input type="checkbox"/>	2.2.5 Shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
K	5, 11	<input type="checkbox"/>	2.2.6 Shellfish are allowed to drain in a clean container or on clean towels prior to opening.
K	2, 5, 11	<input type="checkbox"/>	2.2.7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol, or gloves are donned. The gloves, if worn, are latex, nitrile and/or stainless-steel mesh to protect analyst's hands from injury.
C	5, 11	<input type="checkbox"/>	2.2.8 Shellfish are not shucked through the hinge.
C	5, 11, 12, 15	<input type="checkbox"/>	2.2.9 The contents of the shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
C	12, 15	<input type="checkbox"/>	2.2.10 A representative sample of 10 to 14 shellfish is used for analysis.

C	2, 11	<input type="checkbox"/>	2.2.11 The quantity of meat and liquor is sufficient to cover the blender blades or additional shellfish are used in order to ensure sample homogeneity.
K	5, 12, 13, 15	<input type="checkbox"/>	2.2.12 Either a 1:1 dilution is made, or the sample is homogenized without dilution. If a dilution is made, the sample is weighed to the nearest 0.1 g and an equal amount, by weight, of diluent is added.
K	12, 14, 15	<input type="checkbox"/>	2.2.13 Sterile phosphate buffered saline (pH 7.4) or alkaline peptone water (APW) is used as the sample diluent. If APW is used, sample analysis is conducted immediately.
C	12, 15	<input type="checkbox"/>	2.2.14 Samples are blended at for 90-120 seconds until homogenous.
PART III – ALKALINE PHOSPHATASE PROBE METHOD FOR <i>VIBRIO VULNIFICUS</i> AND <i>VIBRIO PARAHAEMOLYTICUS</i> DETECTION IN SHELLFISH			
			3.1 Preparation of Samples for the Alkaline Phosphatase Probe Method: Direct Plating
C	2, 12, 15	<input type="checkbox"/>	3.1.1 For oyster samples, two tenths (0.20) of a gram of the initial 1:1 diluted homogenate (or 0.10 g of undiluted homogenate) and/or appropriate dilutions are used as inoculum. Dilutions are made in sterile PBS or APW. If APW is used, time from initial dilution until plating does not exceed 30 minutes. For samples other than oysters, 100 µl of the 1:10 dilution and/or subsequent dilutions should be used as inoculum.
K	12, 15	<input type="checkbox"/>	3.1.2 For analysis of total <i>V. parahaemolyticus</i> , at least one (1) T1N3 plate is inoculated to be probed for the <i>tlh</i> gene. For pathogenic <i>V. parahaemolyticus</i> , at least two (2) T1N3 plates are inoculated to be probed for the <i>tdh</i> gene. For analysis of <i>V. vulnificus</i> , at least one (1) VVA plate is inoculated to be probed for the <i>vvhA</i> gene.
K	12, 15	<input type="checkbox"/>	3.1.3 Sterile cell spreaders are used to spread each inoculum evenly onto the dry T1N3 and/or VVA agar plates.
C	2	<input type="checkbox"/>	3.1.4 For <i>V. parahaemolyticus</i> analysis, a <i>tdh</i>+ <i>V. parahaemolyticus</i> culture diluted to <math>10^3</math> per ml is used as a positive process control. A non-<i>V. parahaemolyticus</i> culture is used as a negative process control. For <i>V. vulnificus</i> analysis, a <i>V. vulnificus</i> culture diluted to <math>10^3</math> per ml is used as a positive process control. A non-<i>V. vulnificus</i> culture is used as a negative process control.
C	2	<input type="checkbox"/>	3.1.5 The process control cultures accompany the samples throughout incubation and hybridization and color development phases of the method. Results are recorded, and records are maintained.
C	12, 15	<input type="checkbox"/>	3.1.6 Inoculated plates are incubated 16-24 hours at 35 ± 2 °C. All plates are used for colony lifts and hybridization, except for those with confluent growth.

			3.2 Preparation of Samples for the Alkaline Phosphatase Probe Method: APW Enrichment and Colony Isolation
K	11, 12	<input type="checkbox"/>	3.2.1 Sterile phosphate buffered saline (PBS) is used as the sample diluent.
C	12	<input type="checkbox"/>	3.2.2 The 1:10 dilution is prepared gravimetrically with sterile PBS. All successive dilutions are prepared volumetrically.
C	12, 16	<input type="checkbox"/>	3.2.3 Appropriate sample dilutions are inoculated into sterile APW. Specify dilution(s) used: _____ Specify number of tubes per dilution: _____
C	2	<input type="checkbox"/>	3.2.4 For <i>V. parahaemolyticus</i> analysis, a tdh+ <i>V. parahaemolyticus</i> culture diluted to <math><10^3</math> per ml is used as a positive process control. A non-<i>V. parahaemolyticus</i> culture is used as a negative process control. For <i>V. vulnificus</i> analysis, a <i>V. vulnificus</i> culture diluted to <math><10^3</math> per ml is used as a positive process control. A non-<i>V. vulnificus</i> culture is used as a negative process control.
C	2	<input type="checkbox"/>	3.2.5 The process control cultures accompany the samples throughout incubation, isolation and confirmation. Results are recorded, and records are maintained.
C	12	<input type="checkbox"/>	3.2.6 Inoculated APW enrichment tubes are incubated at 35 ± 2.0 °C.
C	12	<input type="checkbox"/>	3.2.7 Tubes are read after 18-24 hours of incubation. Clear tubes are negative. Turbid tubes are positive. Positive tubes are confirmed as <i>Vibrio parahaemolyticus</i> or <i>Vibrio vulnificus</i> as appropriate.
K	12	<input type="checkbox"/>	3.2.8 A loopful from the top one (1) cm of APW tubes showing growth is streaked onto TCBS for <i>V. parahaemolyticus</i> and mCPC or CC agars for <i>V. vulnificus</i> isolation.
C	12	<input type="checkbox"/>	3.2.9 TCBS plates are incubated at 35 ± 2 °C and mCPC or CC plates are incubated at 35-40 °C for 18-24 hours.
C	12	<input type="checkbox"/>	3.2.10 Presumptive colonies are selected meeting these phenotypic characteristics: a. <i>V. parahaemolyticus</i> appear on TCBS agar as round, opaque, green or bluish colonies, two (2) to three (3) mm in diameter. Interfering large, opaque and yellow colonies are avoided. b. <i>V. vulnificus</i> appear on mCPC or CC agar as round, flat, opaque, yellow colonies, one (1) to two (2) mm in diameter. Typical positives have “fried egg” appearance. Purple/blue colonies are avoided.

C	12	<input type="checkbox"/>	3.2.11 A sterile 96-well microtiter plate is filled with 100 µl/well of APW. Presumptive vibrios are picked from a selective agar plate using a sterile toothpick or wood transfer stick to individual wells. The plate is incubated 3-5 hours or overnight at 35 ± 2 °C. A 48-prong replicator is used to replicate/transfer isolates in the wells to an agar plate (T1N3 for <i>V. parahaemolyticus</i> and VVA for <i>V. vulnificus</i>).
C	12	<input type="checkbox"/>	3.2.12 Plates are incubated at 35 ± 2 °C for 18-24 hours.
3.3 Alkaline Phosphatase Probe Hybridization: Filter Preparation			
C	12, 15	<input type="checkbox"/>	3.3.1 VVA/T1N3 plates are overlaid with labeled (sample number, dilution) #541 Whatman filters for one (1) to 30 minutes.
K	12, 15	<input type="checkbox"/>	3.3.2 Filters are transferred with colony side up to a plastic or glass Petri dish lid containing one (1) ml of lysis solution to wet the filter.
C	12, 15	<input type="checkbox"/>	3.3.3 Filters are microwaved to dryness, but not brown. Microwave for 15-30 seconds/filter, depending on the wattage of the microwave. Additional heating cycles may be required.
K	12, 15	<input type="checkbox"/>	3.3.4 Filters are neutralized for five (5) minutes in an appropriate vessel or container with ammonium acetate (4 ml/filter) on a shaker at room temperature.
C	12, 15	<input type="checkbox"/>	3.3.5 #541 Whatman filters are rinsed two (2) times in 1X SSC buffer (10 ml/filter) for 1-2 minutes. Filters may be air dried and stored at this point.
C	12, 15	<input type="checkbox"/>	3.3.6 Up to 30 filters are incubated in proteinase K solution (10 ml/filter) for 30 minutes at 42 °C with shaking (~50 rpm).
K	12, 15	<input type="checkbox"/>	3.3.7 Filters are rinsed three (3) times in 1X SSC (10 ml/filter) for 10 minutes at room temperature with shaking at 50-125 rpm. Filters may be air dried and stored at this point.
3.4 Alkaline Phosphatase Probe Hybridization: Hybridization.			
C	12, 15	<input type="checkbox"/>	3.4.1 For total <i>V. parahaemolyticus</i> (<i>tlh</i>), the 5'AP-labeled probe 5'aa agc gga tta tgc aga agc act g 3' is used. For pathogenic <i>V. parahaemolyticus</i> (<i>tdh</i>), the 5'AP-labeled probe 5'gg ttc tat tcc aag taa aat gta ttt g 3' is used. For <i>V. vulnificus</i> (<i>vvhA</i>), the 5'AP-labelled probe 5'ga gct gtc acg gca gtt gga acc a 3' is used.
C	12, 15	<input type="checkbox"/>	3.4.2 Probes are stored in the refrigerator and are not frozen.
K	12, 15	<input type="checkbox"/>	3.4.3 A maximum of five (5) filters to be hybridized with the same probe are added to a plastic bag.
C	12, 15	<input type="checkbox"/>	3.4.4 Filters are presoaked in 10-15 ml of hybridization buffer for 30 minutes at 54± 0.1 °C for <i>V. parahaemolyticus</i> (<i>tlh</i> and <i>tdh</i>) or 55 ± 0.1 °C for <i>V. vulnificus</i> with shaking.
C	12, 15	<input type="checkbox"/>	3.4.5 Used buffer is discarded and 10 ml of fresh pre-warmed buffer per bag is added. Probe (final concentration of 0.5 pmol/ml) is quickly added to each bag and incubated for 1 hour at 54 ± 0.1 °C for <i>Vibrio parahaemolyticus</i> or 55 ± 0.1 °C for <i>Vibrio vulnificus</i> with shaking.

K	15	<input type="checkbox"/>	3.4.6 Filters are removed from the bag(s) and transferred to an appropriate vessel or container. Up to 30 filters hybridized with the same probe can be combined.
C	12, 15	<input type="checkbox"/>	3.4.7 Filters are rinsed two (2) times for 10 minutes each in 1X SSC – 1% SDS (for tlh and <i>Vibrio vulnificus</i>) or 3X SSC – 1% SDS (for tdh) (10 ml/filter) at 54 ± 0.1 °C for <i>Vibrio parahaemolyticus</i> or 55 ± 0.1 °C for <i>Vibrio vulnificus</i> with shaking.
K	12, 15	<input type="checkbox"/>	3.4.8 Filters are rinsed five (5) times for five (5) minutes each in 1X SSC (10 ml/filter) at room temperature with shaking.
3.5 Alkaline Phosphatase Probe Hybridization: Color development.			
C	12, 15	<input type="checkbox"/>	3.5.1 In a petri dish containing 20 ml of NBT/BCIP solution, filters (5 or fewer) are added and incubated with gentle shaking at room temperature, or at 35 °C for faster results. The petri dish is kept covered to omit light.
K	12, 15	<input type="checkbox"/>	3.5.2 Color development of the positive control is checked every 30 minutes. Reaction time varies.
K	12, 15	<input type="checkbox"/>	3.5.3 Filters are rinsed in tap or deionized/distilled water (10 ml/filter) three (3) times for 10 minutes each to stop color development.
C	12, 15	<input type="checkbox"/>	3.5.4 Reactions of test sample colonies are compared to the positive and negative process control cultures. Positive reactions appear as purple or brown spots, yellow spots are considered negative reactions. Filters are stored in the dark.
3.6 Alkaline Phosphatase Probe Hybridization: Computation of Results			
C	12, 15	<input type="checkbox"/>	3.6.1 For direct plating, probe-positive colonies are counted and multiplied by the plated dilution factor of the sample to determine the concentration. <u>Note that filter colonies must correspond to colonies visible on the agar plate.</u>
K	15	<input type="checkbox"/>	3.6.2 For direct plating, results are reported as CFU/g of sample.
C	12	<input type="checkbox"/>	3.6.3 For APW enrichment, upon identification of probe-positive colonies refer to the original positive APW dilutions and record MPN value as derived in Appendix 2 of the FDA Bacteriological Analytical Manual (BAM).
K	12, 16	<input type="checkbox"/>	3.6.4 For APW enrichments, results are reported as MPN/g of sample or pass/fail in the case of PHP samples.

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LABORATORY:		DATE OF EVALUATION:	
SHELLFISH LABORATORY EVALUATION CHECKLIST			
SUMMARY OF NONCONFORMITIES			
Page	Item	Observation	Documentation Required

LABORATORY STATUS		
LABORATORY	DATE	
LABORATORY REPRESENTATIVE:		
MICROBIOLOGICAL COMPONENT: (Part I-III)		
A. Results		
Total # of Critical (C) Nonconformities in Parts I-III	_____	
Total # of Key (K) Nonconformities in Parts I-III Total	_____	
# of Critical, Key and Other (O) Nonconformities in Parts I-III	_____	
B. Criteria for Determining Laboratory Status of the Microbiological Component:		
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <p style="margin-left: 40px;">a. The total # of Critical nonconformities is ≥ 4 or</p> <p style="margin-left: 40px;">b. The total # of Key nonconformities is ≥ 13 or</p> <p style="margin-left: 40px;">c. The total # of Critical, Key and Other is ≥ 18</p> <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but ≤ 3.</p>		
C. Laboratory Status (circle appropriate)		
Does Not Conform	Provisionally Conforms	Conforms
<p>Acknowledgment by Laboratory Director/Supervisor:</p> <p>All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before_____.</p> <p>Laboratory Signature: _____ Date:_____</p> <p>LEO Signature: _____ Date:_____</p>		



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Deputy Commissioner

To: ISSC Executive Office
From: J. Michael Hickey, MA DMF, Chief of Shellfish Sanitation and Classification and Susan Boehler, Florence Cenci and Diane Regan, MA DMF, Bacteriologist III
Re: ISSC Request for Interpretation, Date: February 14, 2019
Date: June 18, 2019

Massachusetts Division of Marine Fisheries (MA DMF) would like to comment on the Request for Interpretation received from the ISSC office dated February 14, 2019.

Our review of the interpretation for MO Reference Chapter III, @.01(A) “supporting the NSSP” determined that the MO reference does not address all existing laboratory situations where laboratory support to the NSSP is generated. The interpretation is therefore not complete. The MO reference relies solely on the FDA/State certified Shellfish LEO NSSP technical evaluation to determine laboratory competence and conformance. There are existing laboratory situations however when a FDA/State certified Shellfish LEO NSSP technical evaluation cannot occur. This MO reference therefore must be updated to account for these situations for the benefit of the NSSP and the SSCA. MA DMF has submitted a proposal to the 2019 ISSC conference with a suggested change to this MO reference.

According to the Model Ordinance, a laboratory evaluation conducted by the FDA/State certified Shellfish LEO consists of the following parts:

MO Chapter I @.03 Evaluation of Shellfish Sanitation Program Elements B.1.

“a. Requirements for evaluation of shellfish laboratories shall include at a minimum:

- i. Records audit of laboratory operations both Quality Systems and Technical methods;
- ii. Direct observation of current laboratory operating conditions; and
- iii. Information collection from the Authority and other pertinent sources concerning laboratory operations.”

First a desk audit is made of the laboratories Quality Systems.

MO Chapter 1 @.03 B.1.b.i. Quality System Evaluation. (a)..“All nonconformities must be reconciled prior to scheduling an onsite evaluation of technical methods in NSSP laboratories... The Quality Systems evaluation is performed as a desk audit and is in accordance with the checklist found in Section IV Chapter II.”

Once the laboratory’s Quality System is met with LEO approval, a technical evaluation is conducted onsite using NSSP checklists (MO Section IV. Guidance Documents, Chapter II. Growing Areas, .15 Evaluation of Laboratories by State Shellfish Laboratory Evaluation Officers Including Laboratory Evaluation Checklists). The checklists itemize all technical specifications for the laboratory. The NSSP Laboratory status awarded of Conforms, Provisionally Conforms or Does not Conform is based upon the number and quality of technical violations determined.

The Model Ordinance Chapter 3 @.01C. dictates that a laboratory evaluation will be held at a minimum of every three years as long as the laboratory has satisfactorily passed the initial Quality System Evaluation (QSE). Since no Technical onsite evaluation can occur until the QSE is satisfactorily completed, the time between evaluations could realistically be much longer than 3 years.

MO Chapter 3 @.01C. FDA Responsibilities. "The FDA will ensure that all laboratories generating data in support of the NSSP will be evaluated at a minimum frequency of once every three (3) years."

For NSSP laboratories that have been in operation, inspected previously and had no significant changes this evaluation processes is viable.

The deficiency of the FDA interpretation for MO Chapter III, @.01(A) is that it requires in all cases that a NSSP technical evaluation be performed before a laboratory provides data to "support the NSSP". A technical evaluation however cannot always occur, nor is it always necessary, nor is it the best avenue for assurance to the NSSP that a laboratory's data is validated and legally defensible.

Examples of laboratory situations that are exceptions are:

1. New laboratories to the NSSP
2. NSSP Conforming laboratories performing new NSSP Methods.
3. Laboratories performing Methods that have no Checklists
4. Laboratories that have completed Single Laboratory Validations
5. Laboratories that conform to other externally accredited programs

1. New Laboratories to the NSSP

A new laboratory must first generate a Quality System Manual and have the Quality Systems Evaluation. The laboratory however cannot write a manual until procedures are in place. Laboratory forms, procedures and protocols must be practiced, streamlined and formalized. Once the manual is approved, then a technical evaluation can occur. However, laboratories must be working and generating data for a technical evaluation to take place. As stated in the MO, records are audited, incubator and autoclave charts are examined; the technical evaluation verifies equipment operation under working conditions. It is the direct observation of laboratory operating conditions. According to this interpretation however, the laboratory cannot generate data or operate until an evaluation has occurred. The laboratory cannot have an evaluation until there is data to examine. It is the chicken or the egg situation.

This similar problem occurred for Depuration Plants. Depuration Plants had to operate and generate data to be evaluated, but they could not technically operate until the verification studies were completed. This developed into the Conditional Protocol Verification phase allowing for Depuration plants to support the NSSP while they were being evaluated. New Laboratories should have the same type of interim operational phase that occurs for a time period before an initial laboratory evaluation to allow for similar NSSP support. The FDA/State certified Shellfish LEO should provide cooperative program validation and method support to the new Laboratory until enough data is generated that a formal evaluation can be held. During this Conditional period of intense review, data generation and split sample verification, data output from the laboratory should be used to support the NSSP.

2. NSSP Conforming laboratories performing new NSSP Methods.

A NSSP conforming laboratory may want to introduce a new approved NSSP method in their laboratory such as: mTec, MSC or MPN-Vp. Conforming NSSP Laboratories have already been evaluated by FDA/State certified Shellfish LEO's and have been found to conform to the QSE and to all technical sections for laboratory equipment and methods currently employed. The only subsections that require evaluation are the ones pertaining to the desired new method. In general, the technical aspects of the laboratory have been found to conform to the NSSP. The only question is the ability of the laboratory to produce compliant data with the new method.

A laboratory evaluation demonstrates that the laboratory is technically correct in both equipment and method procedures but successful conformance to a checklist does not measure that the actual outcome of the data produced in the laboratory will be accurate. The NSSP checklist is a living evaluation document; it is not an accreditation or certification document. It is a partnership document between FDA and the states. The checklist is often used for education and training in the state laboratory.

While an evaluation confirms that a laboratory may be technically conforming, the true test of a laboratory's ability to produce data that is standardized within the NSSP occurs with the evaluation of blind proficiency samples. If the laboratory receives unknown spiked samples and can successfully produce data that statistically conforms to other laboratories, using the same methods, then conformance and standardization is actually verified. A NSSP laboratory technical evaluation only confirms that proper conditions are present in the laboratory to hopefully produce conforming data. It is a technical evaluation of conditions. A successful proficiency completion actually verifies that the laboratory and each analyst in that laboratory does produce conforming data. It is a performance evaluation.

There are two commercial performance based proficiencies available to the NSSP laboratory: FDA CFSAN Shellfish and Water Proficiency Test and the NELEOM Performance Evaluation Study. Performance proficiency can also be shown by completing a successfully transferred Method Verification (Section IV, Chapter II. 20) or with split sampling analysis between several laboratories, with at least one laboratory in conforming status for the method.

A FDA/State certified Shellfish LEO evaluation occurs at a minimum of every three years but may be much longer if the QSE is found to have a deficiency on any unrelated issue. If the SSCA/NSSP laboratory has a need to test it usually is in response to an emerging issue, public health threat and/or classification concern. It is to the betterment of the SSCA and therefore to the NSSP that a response with the new method data starts assessing the concern sooner than the three plus year time frame. If a currently conforming Microbiology laboratory can demonstrate performance proficiency in the new microbiology method with a successful performance evaluation, and the verification of that data produced in that laboratory using the NSSP method is confirmed, then that laboratory should be able to support the NSSP with a "conditionally conforming" status. This support will remain in effect until a FDA/State certified Shellfish LEO possessing three years of bench level technical experience in the method is able to conduct the technical evaluation.

3. Laboratories performing Methods that have no Checklists

There are eleven approved methods in the NSSP that do not have checklists: TC-MF mEndo Agar LES, PSP-RBA, DSP- LC-MS/MS, ASP- HPLC, Vv-EIA, Vv-MPN, Vv-QPCR-MPN, Vp-MPN, Vp-PCR, Vp-Direct Plating Method, and Vc-qPCR. In addition there are 5 approved limited use screening methods for Marine Biotxin Testing that do not have checklists.

Most of these non checklist approved methods are for Post Harvest Processing for Vibrios or Marine Biotoxins. For Total Coliform or Vibrio testing, a performance evaluation proficiency test exists: the FDA CFSAN Shellfish and Water Proficiency Test. If a laboratory successfully passes a performance proficiency, should the lack of the existence of a technical checklist stop a NSSP laboratory from supporting the NSSP? Of course not. MA DMF in 1991 completed a performance based SLV for MF mEndo Agar LES. For many years this method was listed in the MO Approved NSSP Test Chart A-10 allowing for its use at MA DMF Newburyport as a regulatory method. Performance proficiency at MA DMF Newburyport is verified yearly by participation in the FDA CFSAN Shellfish and Water Proficiency Test using MF mEndo Agar LES by all analysts in this laboratory. In 2014 a draft checklist of this method was jointly created between FDA and MA DMF. This checklist was recently edited by MA DMF and was submitted as a proposal at the 2019 ISSC conference to assist the NSSP in the method's use. If a laboratory has proven its ability by completing a successful performance evaluation for a method then the laboratory should be awarded "conditionally conforming" status and continue to support the NSSP with data from that method.

4. Laboratories that have completed Single Laboratory Validations (SLV)

Any new laboratory method for consideration in the NSSP must submit a SLV to the ISSC Laboratory Committee. The ISSC Laboratory Committee reviews the extensive SLV critique of method data and if found suitable the package is submitted to the conference for adoption as a NSSP method. Eight SLV submissions have become NSSP Methods and are now listed in the current MO Section IV Guidance Documents – Chapter II. Growing Areas, .14 Approved NSSP Laboratory Tests. Laboratories that completed SLV's with the dates completed are the Laboratories of: Dr. Anita Wright 2009 , Dr. Fran Van Dohl, NOAA 2013; Dr. Jon Deeds FDA 2017; FDA Dauphin Island 2015; Abraxis LLC 2009; Jellett Rapid Testing Ltd, 2005; Neogen Corp, 2013 and Marbionc Dev. Group LLC 2017. The submission of data in a Single Laboratory Validation for the potential adoption of a new NSSP method clearly supports the NSSP. Were these laboratories in the status of NSSP Conforming or Provisionally Conforming at this time of their SLV submission? A SLV is a method performance evaluation performed at a laboratory. Once a method performance is validated, the method's technical aspects are determined with the drafting of a required technical checklist. Under the current FDA interpretation, an SLV with accompanying technical checklist could be accepted as an approved NSSP method by the conference, yet the submitting laboratory would not be able to "support the NSSP" with data from their own method. The FDA interpretation indicates that a FDA Shellfish LEO, who is unfamiliar with the technical or performance of the method, (it could not be a FDA certified State LEO since they do not have three years of bench experience with the SLV method) would have to conduct a technical evaluation at the SLV's laboratory using the SLV laboratory's own devised checklist in order for that lab to be "conforming" and able to start generating data in "support of the NSSP". This is not logical. If a laboratory has proven its ability to perform a method successfully with statistically validated, legally defensive data, then the laboratory should be awarded "conditionally conforming" status to support the NSSP with data.

5. Laboratories that conform to other external accredited programs

The NSSP checklists recognize and contain references to other certified laboratory programs such as the US FDA Milk Program, US EPA drinking water/wastewater and accreditation programs such as ISO and AOAC. Laboratories that conform to these other program certification and accreditation standards are required to submit to annual proficiency sampling for their accredited program's standardization. The data from these laboratories therefore has been validated and found compliant for their program use. If the SSCA is using validated data supplied by an external accredited/certified laboratory for the dual purpose of the protection of shellfish resources and public health, the NSSP should continue to recognize and acknowledge the external accreditation/certification process required of these other programs that

the NSSP itself has referenced. Laboratories with these accreditations and certifications have usually complemented or exceeded the requirements of the evaluation process of the NSSP and should be awarded “conditionally conforming” status to support the NSSP.

The FDA/State certified Shellfish LEO evaluation is an important component of the laboratory evaluation process but it is not the only component. Technical competence must be combined with performance competence for the true standardization of laboratories and output data within the NSSP laboratory. Words without action do not verify results.

As a simple analogy: a technical evaluation results in two states A and B having technically conforming cars. Both have engines, horns, lights, tires, seats; they are technically equal and conform to the same standards. But what if state A car was driven by a race car driver and state B was driven by a 5 year old? What if the state A car was a Jaguar and the state B car was a Honda? Technically they are both cars, but would they drive the same? Would they cross the finish line similarly ending at the same spot? The NSSP should want to know that all labs and all analysts driving conforming cars are crossing the finish line at the same spot. Is the count of 14 FC/100ml from NSSP state A lab the same as a count of 14 FC/100ml from NSSP state B lab? The labs are technically the same (conforming) but with the two different drivers the outcomes could be very different. If both labs are able to produce validated counts of 14 FC/100ml, (8 times in a row, no less) in the absence of a technical evaluation, the implication can be made that the laboratory is operating technically correct with no detrimental conditions that are impacting results since compliant data was produced. If a lab is not able to validate counts within a proficiency this should prompt a NSSP technical evaluation to determine what technical conditions are occurring which are detrimental to the laboratory’s results.

Completion of a technical evaluation validates and standardizes equipment and technical specifications in the NSSP laboratory but only implies that laboratory performance may produce conforming data. Technical evaluations occur every 3-5 years. Successful completion of a performance evaluation validates and confirms the standardization of data in the NSSP laboratory and implies technical conformance. Performance evaluations occur yearly allowing for better surveillance. The ability of a laboratory to produce statistically confirmed standardized data is the primary goal of the NSSP, not just to be technically correct. Technical evaluations are subject to human interpretation as LEO’s must account for differing conditions and equipment. Performance evaluations rely on statistics. The completion of a performance evaluation provides greater assurance than the technical evaluation and is therefore a stronger legally defensible procedure and document. An additional laboratory category of “conditionally conforming” should be included in the MO to allow for laboratories that have completed a performance evaluation to support the NSSP with validated data.

The primary goal of the NSSP, FDA and the SSCA is to assure that laboratories are producing verified and accurate data. The technical evaluation assures that conditions are present in that laboratory for successful data to be produced; it is proactive. It does not however assure that the production of successful data actually occurs nor can the technical evaluation be used for all laboratory situations which support the NSSP. The MA DMF response has outlined 5 instances where technical evaluations cannot occur but where laboratories are able to contribute validated, legally defensible data “supporting” the SSCA and the NSSP; with the cooperative goal of promoting the best interests of the shellfish industry and public health.

MA DMF appreciates the opportunity to comment. MA DMF is available for further discussion and questions.

Attachment: MA DMF 2019 Proposal submission “Conditionally Conforming Laboratory Status”

RHODE ISLAND DEPARTMENT OF HEALTH - OFFICE OF FOOD PROTECTION,
RM 203, CANNON BLDG., 3 CAPITOL HILL, PROVIDENCE, RI 02908

LAB #

SAMPLE COLLECTION FORM

210957 12 AM 10:09

DATE OF COLLECTION 12/12/01	DATE OF ANALYSIS 12/20-21/01	DATE OF REPORT 12/14/01	LAB SUP PCE
CONDITION HOT FROZEN YES COLD OTHER NO	ITEM OYSTER'S	PRODUCT CODE/DOM	MOP
SEALING SEALING YES NO	BRAND NAME Geib	TEMP ESTAB	TEMP REC °C
EXP DATE	SIZE	TEMP ANALYSIS	REASON 01
DATE OF SHIP	FROM LOT OF	COLLECTED AT Little Allen's HARBOR	REASON
SHIPPER/PACKER & ADDRESS 76 Quaker Dr West Warwick 02883		FOLLOW UP SAMPLE #1	SOURCE 10
COMPLAINT*		ORIGINAL CONTAINER YES NO	DATE PURCHASED 12/13/01
NAME & LOCATION OF STORE WHERE PURCHASED		PRODUCT USED YES NO	AMOUNT REMARKS
HOW STORED FROZEN COLD	AMBIENT	IMPORT PRODUCT YES NO	INTERVIEWED BY
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:			

SIGNED FROM John Meller	SIGNED TO [Signature]	DATE 12-12-01	TIME 10:15 AM	HOW STORED in ice
AGENCY # 164 John Meller	PRINT NAME THEODORE L. LUKAS			
SIGNED FROM [Signature]	SIGNED TO			
PRINT NAME THEODORE L. LUKAS	PRINT NAME	12-12-01	10:30 AM	

1. CEREAL	2. MEAT PROT	3. TOT H ₂ O	4. SOY FL	5. NFDN	6. ADD H ₂ O	7. TOT PROT	8. TOT FAT
9. SORBATES	10. TBA	11. FFA	12. WT/VOL	13. NaNO ₂	14. EL'TROPHO	15. TOTAL SOLIDS	16. HISTAMINE
17. NaCl	18. NH ₃	19. SULFITE	20. Hg	21. TVB	22. DOMOIC ACID	23. TMA	24. VITAMIN A
25. pH	26. BRIX	27. INSECT	28. RODENT	29. BENZOT'S	30. ASCORB'S	31. INDOLE	32. VITAMIN D
33. PKG INT	34. AFLATOXIN	35. Pb 4.34	36. Cd 0.53	37. Cu 61.7	38. Cr 0.98	39. Zn 385	40. A _w
41. Na	42. TSP	43. Ca	44. P	45. Mg	46. GLUTAMIC/MSG	47. ACETIC ACID	48. GLUCOSE
47. FRUCTOSE	48. SUCROSE	49. LACTOSE	50. MALTOSE	51. YEAST	52. MOLD	53. PSP	54. OTHER
56. FECAL/MPN	57. TOT COLIFORM/MPN	58. CL BOT	59. CL PERF	60. COAG STAPH	61. NON-COAG STAPH	62. LISTERIA	63. CAMPYLO
64. SPC	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC			

COLLECTOR'S NOTES:
*Samples from upwells
Bill Geib lease
WATER Temp 43°*

LABORATORY NOTES:
*Ni: 1.86 ppm
Pb: 244 ppm*

RECEIVED

JAN - 2 2002

RI DEPARTMENT OF HEALTH
DIVISION OF FOOD PROTECTION

EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)	ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE	CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY	REVIEWED BY:
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SAMPLE COLLECTION FORM

DATE OF COLLECTION 3/28/02 <i>Paul By</i>		DATE OF ANALYSIS 3-29-02	DATE OF REPORT 4.2.02	LAB SUP PLP
CONDITION HOT FROZEN COLD OTHER	SEALED YES NO	ITEM OYSTERS		PRODUCT CODE/DOM
EXP DATE	SIZE	BRAND NAME		MOP
DATE OF SHIP	FROM LOT OF	COLLECTED AT Geib lease		REASON 01
SHIPPER/PACKER & ADDRESS				TEMP ESTAB
				TEMP REC °C
				TEMP ANALYSIS
				FOLLOW UP SAMPLE #1 SM 11 SM 13 SM 17
				SOURCE 10
				ANALYSIS 246,13
				DATE PURCHASED
				CONTAINER YES NO

COMPLAINT*				PRODUCT USED YES NO
NAME & LOCATION OF STORE WHERE PURCHASED				AMOUNT REMAINING
HOW STORED FROZEN COLD	AMBIENT	IMPORT PRODUCT YES NO	INTERVIEWED BY	DATE
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:				TIME

SIGNED FROM <i>John Miller</i>	SIGNED TO <i>Phung Tam</i>	DATE 3-29-02	TIME 9:30	HOW STORED in cooler
AGENT PRINT NAME 164 John Miller	PRINT NAME			
SIGNED FROM <i>Phung Tam</i>	SIGNED TO <i>Kerry Patterson</i>	DATE 3-29-02	TIME 10:00	HOW STORED in refrigerator
PRINT NAME Phung Tam	PRINT NAME Kerry Patterson			
SIGNED FROM <i>Kerry Patterson</i>	SIGNED TO <i>Chris Edley</i>	DATE 4/2/02	TIME 10:40	HOW STORED REF
PRINT NAME Kerry Patterson	PRINT NAME Chris Edley			

1. CEREAL	2. MEAT PROT	3. TOT H ₂ O	4. SOY FL	5. NFDM	6. ADD H ₂ O	7. TOT PROT	8. TOT FAT
%	%	%	%	%	%	%	%
9. SORBATES	10. TBA	11. FFA	12. WT/VOL	13. NaNO ₂	14. EL'TROPHO	15. TOTAL SOLIDS	16. HISTAMINE
ppm	ug ml/g	%	%	ppm	ppm	%	mg/100g
17. NaCl	18. NH ₃	19. SULFITE	20. Hg	21. TVB	22. DOMOIC ACID	23. TMA	24. VITAMIN A
%	mg/100g	ppm	ppm	mg N/100g	ug/ml	mgN/100g	IU/Qt
25. pH	26. BRIX	27. INSECT	28. RODENT	29. BENZOT'S	30. ASCORB'S	31. INDOLE	32. VITAMIN D
	%			ppm	ppm	ug/100g	IU/Qt
33. PKG INT	34. AFLATOXIN	35. Pb	36. Cd	37. Cu	38. Cr	39. Zn	40. A _w
	ppb	ppm	ppm	ppm	ppm	ppm	
41. Na	42. TSP	43. Ca	44. P	45. Mg	46. GLUTAMIC/MSG	47. ACETIC ACID	48. GLUCOSE
mg/Serving		mg/100g	mg/100g	mg/100g	%	mg/100g	%
47. FRUCTOSE	48. SUCROSE	49. LACTOSE	50. MALTOSE	51. YEAST	52. MOLD	53. PSP	54. OTHER
%	%	%	%	%	%	%	%
56. FECAL/MPN 220	57. TOT COLIFORM/MPN 220	58. CL BOT	59. CL PERF	60. COAG STAPH	61. NON-COAG STAPH	62. LISTERIA	63. CAMPYLO
64. SFC 730	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC			

COLLECTOR'S NOTES:
*Samples from grow out area. Placed there 9901
 Grown AT lease only*

LABORATORY NOTES:
Fe: 220 ppm
Ni: 1.32 ppm

EVALUATION BY LAB:	ADMINISTRATION FOLLOW-UP:	CONCLUSION:	REVIEWED BY:
1. NOT AN FP OBLIGATION	1. CHECK ESTABLISHMENT	1. SATISFACTORY	
2. FP VIOLATION - NO VIOLATION	2. ADDITIONAL SAMPLES	2. QUESTIONABLE	
3. FP ACTION INDICATED	3. EMBARGO	3. VIOLATES STD	
4. INSUFFICIENT SAMPLE FOR ANALYSIS	4. REVIEW PROCESS	4. UNSATISFACTORY	
5. INSUFFICIENT INFORMATION FOR ANALYSIS	5. NO FURTHER ACTION		
6. IMPROPER CHAIN OF CUSTODY	6. DISPOSE		
7. LAB UNABLE TO PERFORM TESTING (REASON)			

RHODE ISLAND DEPARTMENT OF HEALTH - OFFICE OF FOOD PROTECTION,
RM 203, CANNON BLDG., 3 CAPITOL HILL, PROVIDENCE, RI 02908

LAB # **14**
11/15 29 AM 9:22

SAMPLE COLLECTION FORM

DATE OF COLLECTION 3/28/02		DATE OF ANALYSIS 3-25-02	DATE OF REPORT 4-2-02	LAB SUP PLJ	SHIPPER/PACKER & ADDRESS		
CONDITION HOT FROZEN YES COLD OTHER NO		ITEM OYSTERS		PRODUCT CODE/DOM	MOP	SAMPLE CODE 61	
SEALING YES NO		BRAND NAME		TEMP ESTAB	TEMP REC °C	TEMP ANALYSIS	
DATE OF SHIP		FROM LOT OF		COLLECTED AT Geib lease		REASON 01	
SHIPPING/PAKING		ADDRESS		FOLLOW UP SAMPLE #1 SM 11 SM 13 SM 24		SOURCE 10	
COMPLAINT*				ORIGINAL CONTAINER YES NO		DATE PURCHASED 29(8.13)	
NAME & LOCATION OF STORE WHERE PURCHASED				PRODUCT USED YES NO			
HOW STORED FROZEN COLD		IMPORT PRODUCT YES NO		INTERVIEWED BY	DATE	TIME	AMOUNT REMAINING
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC...							
SIGNED FROM John Miller		SIGNED TO Phung Tam		DATE 3-29-02	TIME 9:30	HOW STORED in cooler	
AGENT PRINT NAME John Miller		PRINT NAME Phung Tam					
SIGNED FROM Phung Tam		SIGNED TO Kerry Patterson		DATE 3-29-02	TIME 10:00	in Refrigerator	
PRINT NAME Phung Tam		PRINT NAME Kerry Patterson					
SIGNED FROM Kerry Patterson		SIGNED TO Chris Ely		DATE 4/2/02	TIME 10:40	REF	
PRINT NAME BERRY PATTERSON		PRINT NAME Chris Ely					
1. CEREAL %	2. NEXT PROT %	3. TOT H ₂ O %	4. SOY FL %	5. NFDN %	6. ADD H ₂ O %	7. TOT PROT %	8. TOT FAT %
9. SORBATES ppm	10. TBA ug mal/g	11. FFA %	12. WT/VOL %	13. NaNO ₂ ppm	14. EL'TROPHO ppm	15. TOTAL SOLIDS %	16. HISTAMINE mg/100g
17. NaCl %	18. NH ₃ mg/100g	19. SULFITE ppm	20. Hg ppm	21. TVB mg N/100g	22. DOMOIC ACID ug/ml	23. TMA mgN/100g	24. VITAMIN A IU/Oz
25. pH	26. BRUX %	27. INSECT %	28. RODENT %	29. BENZOT'S ppm	30. ASCORB'S ppm	31. INDOLE ug/100g	32. VITAMIN D IU/Oz
33. PKG INT	34. AFLATOXIN ppb	35. Pb 0.20 ppm	36. Cd 0.26 ppm	37. Cu 27.3 ppm	38. Cr 0.37 ppm	39. Zn 426 ppm	40. A _w
41. Na mg/Serving	42. TSP mg/100g	43. Ca mg/100g	44. P mg/100g	45. Mg mg/100g	46. GLUTAMIC/MSG %	47. ACETIC ACID mg/100g	48. GLUCOSE %
47. FRUCTOSE %	48. SUCROSE %	49. LACTOSE %	50. MALTOSE %	51. YEAST %	52. MOLD %	53. PSP %	54. OTHER %
56. FECAL/MPN 20	57. TOT COLIFORM/MPN 50	58. CL BOT	59. CL PERF	60. COAG STAPH	61. NON-COAG STAPH	62. LISTERIA	63. CAMPYLO
64. SPC 770	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC			
COLLECTOR'S NOTES: <i>Sample from grow out Placed thru SUMMER 2000 2 WITNESS AT LEASE</i>				LABORATORY NOTES: LAB 14 Fe: 21.4 ppm Ni: 1.24 ppm			
EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)				ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE		CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY	
FOOD QUALITY (SAFETY SHELF-LIFE & CONSUMER ACCEPTANCE) INVOLVES THE ENTIRE CHAIN OF PRODUCTION PROCESSING & DISTRIBUTION OF FOOD*				REVIEWED BY:			
* SAMPLE WILL BE MAINTAINED FOR THIRTY (30) DAYS AFTER ANALYSIS AND THEN DISPOSED OF BY THE LABORATORY.				Supporting Documentation Task Force I & Task Force II - Page 297			
WHITE - DATA ENTRY		YELLOW - LAB		PINK - FOOD PROTECTION		GOLDENROD - CONSUMER	

LAB # **13**

SAMPLE COLLECTION FORM

213114 25 AM 9:22

DATE OF COLLECTION 3/29/02		DATE OF ANALYSIS 3.29.02	DATE OF REPORT 4.2.02	LAB SUR PCE
CONDITION HOT FROZEN COLD OTHER	SEALED YES NO	ITEM OYSTERS		PRODUCT CODE/DOM
EXP DATE	SIZE	BRAND NAME	TEMP ESTAB	TEMP REC °C
DATE OF SHIP	FROM LOT OF	COLLECTED AT Geib lease	REASON 01	
SHIPPER/PACKER & ADDRESS			ADDRESS	SOURCE 10
COMPLAINT*			DATE PURCHASED	ANALYSIS 24(8,13)
NAME & LOCATION OF STORE WHERE PURCHASED			ORIGINAL CONTAINER YES NO	PRODUCT USED YES NO
HOW STORED FROZEN COLD	AMBIENT	IMPORT PRODUCT YES NO	INTERVIEWED BY	DATE
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:				
SIGNED FROM John Miller	AGENT # 164 John Miller	SIGNED TO Phan Tu	DATE 3-29-02	TIME 9:30
PRINT NAME John Miller		PRINT NAME Phan Tu	HOW STORED in cooler	
SIGNED FROM Phan Tu	AGENT #	SIGNED TO Kerry Peterson	DATE 3-29-02	TIME 10:00
PRINT NAME Phan Tu		PRINT NAME Kerry Peterson	HOW STORED in refrigerator	
SIGNED FROM Kerry Peterson	AGENT #	SIGNED TO Chris Ellis	DATE 4/6/02	TIME 10:45
PRINT NAME Kerry Peterson		PRINT NAME Chris Ellis	HOW STORED ref	
1. CEREAL	2. MEAT PROT	3. TOT H ₂ O	4. SOY FL	5. NFDM
%	%	%	%	%
9. SORBATES	10. TBA	11. FFA	12. WT/VOL	13. NaNO ₂
ppm	ug ml/g	%	%	ppm
17. NaCl	18. NH ₃	19. SULFITE	20. Hg	21. TVB
%	mg/100g	ppm	ppm	mg N/100g
25. pH	26. BRIX	27. INSECT	28. RODENT	29. BENZO'S
		%		ppm
33. PKG INT	34. AFLATOXIN	35. Pb	36. Cd	37. Cu
	ppb	ppm	ppm	ppm
41. Na	42. TSP	43. Ca	44. P	45. Mg
mg/Serving		mg/100g	mg/100g	mg/100g
47. FRUCTOSE	48. SUCROSE	49. LACTOSE	50. MALTOSE	51. YEAST
%	%	%	%	mg/100g
56. FECAL/MPN <20	57. TOT COLIFORM/MPN <20	58. CL BOT	59. CL PERF	60. COAG STAPH
				61. NON-COAG STAPH
64. SFC 2100	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC

COLLECTOR'S NOTES:
*Sample from grow out
 Placed there Summer June-Oct. 2001*

LABORATORY NOTES:
LAB# 13
Fe: 28.1 ppm
Ni: 620 ppm

EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)	ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE	CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY	REVIEWED BY:
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SAMPLE COLLECTION FORM

DATE OF COLLECTION 8/4/98		Ann BX NCS		DATE OF ANALYSIS 8/13/98	DATE OF REPORT 8/17/98	LAB SUP PCE			
CONDITION HOT FROZEN COLD OTHER	SCALED YES NO	ITEM OYSTERS			PRODUCT CODE/DOM	MOP	SAMPLE CODE 61		
EXP DATE	SIZE	BRAND NAME			TEMP ESTAB	TEMP REC °C	TEMP ANALYSIS		
DATE OF SHIP	FROM LOT OF	COLLECTED AT Billington Cove Marina				REASON 01			
SHIPPER/PACKER & ADDRESS						FOLLOW UP SAMPLE #1		SOURCE 10	
COMPLAINT*						ORIGINAL CONTAINER YES NO		ANALYSIS 13	
NAME & LOCATION OF STORE WHERE PURCHASED						DATE PURCHASED		PRODUCT USED YES NO	
HOW STORED FROZEN COLD	AMBIENT	IMPORT PRODUCT YES NO	INTERVIEWED BY	DATE	TIME	AMOUNT REMAINING			
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:									
SIGNED FROM <i>[Signature]</i>		SIGNED TO <i>[Signature]</i>		DATE 8/11/98	TIME 11:40	HOW STORED Ambient			
AGENT'S PRINT NAME 164 John Mullen		PRINT NAME MULLER SILVIA							
SIGNED FROM		SIGNED TO							
PRINT NAME		PRINT NAME							
SIGNED FROM		SIGNED TO							
PRINT NAME		PRINT NAME							
1. CEREAL	2. MEAT PROT	3. TOT H ₂ O	4. SOY FL	5. NFDN	6. ADD H ₂ O	7. TOT PROT	8. TOT FAT		
%	%	%	%	%	%	%	%		
9. SORBATES	10. TBA	11. FFA	12. WT/VOL	13. NaNO ₂	14. EL'TROPHO	15. TOTAL SOLIDS	16. HISTAMINE		
ppm	ug mal/g	%	%	ppm	ppm	%	mg/100g		
17. NaCl	18. NH ₃	19. SULFITE	20. Hg	21. TVB	22. DOMOIC ACID	23. TMA	24. VITAMIN A		
%	mg/100g	ppm	ppm	mg N/100g	ug/ml	mgN/100g	IU/Qt		
25. pH	26. BRIX	27. INSECT	28. RODENT	29. BENZOT'S	30. ASCORB'S	31. INDOLE	32. VITAMIN D		
	%	%	%	ppm	ppm	ug/100g	IU/Qt		
33. PKG INT	34. AFLATOXIN	35. Pb	36. Cd	37. Cu	38. Cr	39. Zn	40. A _w		
	ppb	0.47 ppm	0.45 ppm	127 ppm	0.15 ppm	798 ppm			
41. Na	42. TSP	43. Ca	44. P	45. Mg	46. GLUTAMIC.MSG	47. ACETIC ACID	48. GLUCOSE		
mg/Serving		mg/100g	mg/100g	mg/100g	%	mg/100g	%		
47. FRUCTOSE	48. SUCROSE	49. LACTOSE	50. MALTOSE	51. YEAST	52. MOLD	53. PSP	54. OTHER		
%	%	%	%	%	%	%			
56. FECAL/MPN	57. TOT COLIFORM/MPN	58. CL BOT	59. CL PERF	60. COAG STAPH	61. NON-COAG STAPH	62. LISTERIA	63. CAMPYLO		
64. SPC	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC					
COLLECTOR'S NOTES: <i>Moanston Oyster nursery stock. appx. 36mm</i>				LABORATORY NOTES: <i>Fe: 133 ppm</i>					
EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)				ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE			CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY		REVIEWED BY:
FOOD QUALITY (SAFETY SHELF-LIFE & CONSUMER ACCEPTANCE) INVOLVES THE ENTIRE CHAIN OF PRODUCTION PROCESSING & DISTRIBUTION OF FOOD*									
* SAMPLE WILL BE MAINTAINED FOR THIRTY (30) DAYS AFTER ANALYSIS AND THEN DISPOSED OF BY THE LABORATORY.									
WHITE - DATA ENTRY YELLOW - Supporting Documentation Task Force GOLDENROD - CONSUMER									

SAMPLE COLLECTION FORM

202209 16 PM 1:52

DATE OF COLLECTION 11/29/00		DATE OF ANALYSIS 11/29/00	DATE OF REPORT 11/30/00	LAB SUP POE			
CONDITION HOT FROZEN COLD OTHER	SEALED YES NO	ITEM OYSTERS	PRODUCT CODE/DOM	MOP	SAMPLE CODE 61		
EXP DATE	SIZE	BRAND NAME	TEMP ESTAB	TEMP REC °C	TEMP ANALYSIS		
DATE OF SHIP	FROM LOT OF	COLLECTED AT Point Judith Pond	ADDRESS Beach Island		FOLLOW UP SAMPLE #1	REASON 04	
SHIPPER/PACKER & ADDRESS					ANALYSIS 13	SOURCE 10	
COMPLAINT*					ORIGINAL CONTAINER YES NO	DATE PURCHASED	
NAME & LOCATION OF STORE WHERE PURCHASED					PRODUCT USED YES NO		
HOW STORED FROZEN COLD	AMBIENT	IMPORT PRODUCT YES NO	INTERVIEWED BY	DATE	TIME	AMOUNT REMAINING	
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:							
SIGNED FROM John Muller	SIGNED TO Chris Ellis	DATE 11/16/00	TIME 2:00	HOW STORED on ice			
AGENT #	PRINT NAME 169 John Muller	PRINT NAME Chris Ellis					
SIGNED FROM	SIGNED TO	PRINT NAME					
SIGNED FROM	SIGNED TO	PRINT NAME					
SIGNED FROM	SIGNED TO	PRINT NAME					
1. CEREAL	2. MEAT PROT	3. TOT H ₂ O	4. SOY FL	5. NFDM	6. ADD H ₂ O	7. TOT PROT	8. TOT FAT
%	%	%	%	%	%	%	%
9. SORBATES	10. TBA	11. FFA	12. WT/VOL	13. NaNO ₂	14. EL'TROPHO	15. TOTAL SOLIDS	16. HISTAMINE
ppm	ug mal/g	%		ppm		%	mg/100g
17. NaCl	18. NH ₃	19. SULFITE	20. Hg	21. TVB	22. DOMOIC ACID	23. TMA	24. VITAMIN A
%	mg/100g	ppm	ppm	mg N/100g	ug/ml	mgN/100g	IU/Ot
25. pH	26. BRIX	27. INSECT	28. RODENT	29. BENZOT'S	30. ASCORB'S	31. INDOLE	32. VITAMIN D
	%	%	%	ppm	ppm	ug/100g	IU/Ot
33. PKG INT	34. AFLATOXIN	35. Pb	36. Cd	37. Cu	38. Cr	39. Zn	40. A _w
	ppb	0.77 ppm	0.17 ppm	25.4 ppm	0.34 ppm	590 ppm	
41. Na	42. TSP	43. Ca	44. P	45. Mg	46. GLUTAMIC/MSG	47. ACETIC ACID	48. GLUCOSE
mg/Serving		mg/100g	mg/100g	mg/100g	%	mg/100g	%
47. FRUCTOSE	48. SUCROSE	49. LACTOSE	50. MALTOSE	51. YEAST	52. MOLD	53. PSP	54. OTHER
%	%	%	%	%	%	%	%
56. FECAL/MPN	57. TOT COLIFORM/MPN	58. CL BOT	59. CL PERF	60. COAG STAPH	61. NON-COAG STAPH	62. LISTERIA	63. CAMPYLO
64. SPC	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC			
COLLECTOR'S NOTES: Samples collected by Bot R results				LABORATORY NOTES: Fe: 26.9 ppm			
EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)				ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE		CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY	
						REVIEWED BY:	
FOOD QUALITY (SAFETY SHELF-LIFE & CONSUMER ACCEPTANCE) INVOLVES THE ENTIRE CHAIN OF PRODUCTION PROCESSING & DISTRIBUTION OF FOOD*							
* SAMPLE WILL BE MAINTAINED FOR THIRTY (30) DAYS AFTER ANALYSIS AND THEN DISPOSED OF BY THE LABORATORY.							
WHITE - DATA ENTRY							

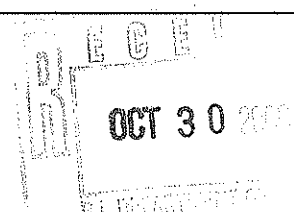
RHODE ISLAND DEPARTMENT OF HEALTH - OFFICE OF FOOD PROTECTION,
RM 203, CANNON BLDG., 3 CAPITOL HILL, PROVIDENCE, RI 02908

LAB #

SAMPLE COLLECTION FORM

198251 13 AM 10:32

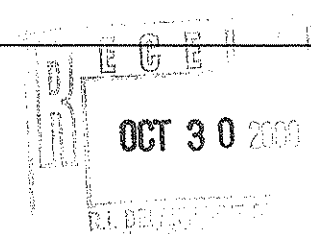
DATE OF COLLECTION <i>7/13/00</i>		DATE OF ANALYSIS <i>10/27/00</i>		DATE OF REPORT <i>10/27/00</i>		LAB SUP <i>PCE</i>	
CONDITION HOT FROZEN COLD OTHER		ITEM <i>OYSTERS</i>		PRODUCT CODE/DOM		MOP	
SIGNED <i>YES</i>		BRAND NAME		TEMP ESTAB		TEMP REC °C	
EXP DATE		SIZE		TEMP ANALYSIS <i>66</i>		REASON <i>01</i>	
DATE OF SHIP		FROM LOT OF		COLLECTED AT <i>MOONSTONE OYSTER</i>		SOURCE <i>02</i>	
SHIPPER/PACKER & ADDRESS		ADDRESS		FOLLOW UP SAMPLE #1		ANALYSIS <i>13</i>	
COMPLAINT*						ORIGINAL CONTAINER YES NO	DATE PURCHASED
NAME & LOCATION OF STORE WHERE PURCHASED						PRODUCT USED YES NO	AMOUNT REMAINING
HOW STORED FROZEN SOLD		AMBIENT		IMPORT PRODUCT YES NO		INTERVIEWED BY	
DATE		TIME		DATE		TIME	
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:							
SIGNED FROM AGENT # PRINT NAME		SIGNED TO PRINT NAME		DATE <i>7/13/00</i>		TIME <i>10:35</i>	
SIGNED FROM		SIGNED TO		DATE		TIME	
PRINT NAME SIGNED FROM		PRINT NAME SIGNED TO		DATE		TIME	
PRINT NAME		PRINT NAME		DATE		TIME	
1. CEREAL		2. MEAT PROT		3. TOT H ₂ O		4. SOY FL	
%		%		%		%	
9. SORBATES		10. TBA		11. FFA		12. WT/VOL	
ppm		ug mal/g		%		%	
17. NaCl		18. NH ₃		19. SULFITE		20. Hg	
%		mg/100g		ppm		ppm	
25. pH		26. BRIX		27. INSECT		28. RODENT	
		%		%		%	
33. PKG INT		34. AFLATOXIN		35. Pb		36. Cd	
		ppb		<i><0.54</i> ppm		<i>0.19</i> ppm	
41. Na		42. TSP		43. Ca		44. P	
mg/Serving		mg/100g		mg/100g		mg/100g	
47. FRUCTOSE		48. SUCROSE		49. LACTOSE		50. MALTOSE	
%		%		%		%	
56. FECAL/MPN		57. TOT COLIFORM/MPN		58. CL BOT		59. CL PERF	
		%		%		%	
64. SPC		65. SALMONELLA		66. B CEREUS		67. YERSINIA	
						ORGANOLEPTIC	
COLLECTOR'S NOTES: <i>Cage #24 oldest system on lease placed mid-late 98</i>				LABORATORY NOTES: <i>Fe: 31.4 ppm</i>			
EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)				ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE		CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY	
FOOD QUALITY (SAFETY SHELF-LIFE & CONSUMER ACCEPTANCE) INVOLVES THE ENTIRE CHAIN OF PRODUCTION PROCESSING & DISTRIBUTION OF FOOD*				REVIEWED BY:			
* SAMPLE WILL BE MAINTAINED FOR THIRTY (30) DAYS AFTER ANALYSIS AND THEN DISPOSED OF BY THE LABORATORY.				Supporting Documentation Task Force I & Task Force II - Page 301			
WHITE - DATA ENTRY				GOLDENROD - CONSUMER			



SAMPLE COLLECTION FORM

198250 13 AM 10:32

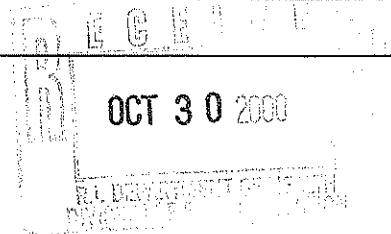
DATE OF COLLECTION 7/13/00		DATE OF ANALYSIS 10/12/00		DATE OF REPORT 10/12/00		LAB SUP PCE	
CONDITION HOT FROZEN COLD OTHER		ITEM OYSTERS		PRODUCT CODE/DOM		MOP	
SEALING YES NO		BRAND NAME OYSTERS		TEMP ESTAB		TEMP REC °C	
EXP DATE		SIZE		TEMP ANALYSIS		SAMPLE CODE 661	
DATE OF SHIP		FROM LOT OF		COLLECTED AT MOONSTONE OYSTER		REASON 01	
SHIPPER/PACKER & ADDRESS		ADDRESS		FOLLOW UP SAMPLE #1		SOURCE 02	
COMPLAINT*		ORIGINAL CONTAINER YES NO		DATE PURCHASED		ANALYSIS 13	
NAME & LOCATION OF STORE WHERE PURCHASED		PRODUCT USED YES NO		AMOUNT REMAINING			
HOW STORED FROZEN COLD		AMBIENT		IMPORT PRODUCT YES NO		INTERVIEWED BY	
DATE		TIME		TIME		AMOUNT REMAINING	
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.:							
SIGNED FROM John Miller		SIGNED TO Tracey Lempelton		DATE 7/13/00		TIME 10:35	
AGENT PRINT NAME 164 John Miller		PRINT NAME TRACEY LEMPELTON		HOW STORED ambient			
SIGNED FROM		SIGNED TO					
PRINT NAME		PRINT NAME					
SIGNED FROM		SIGNED TO					
PRINT NAME		PRINT NAME					
1. CEREAL	2. MEAT PROT	3. TOT H ₂ O	4. SOY FL	5. NFDM	6. ADD H ₂ O	7. TOT PROT	8. TOT FAT
%	%	%	%	%	%	%	%
9. SORBATES	10. TBA	11. PFA	12. WT/VOL	13. NaNO ₂	14. EL'TROPHO	15. TOTAL SOLIDS	16. HISTAMINE
ppm	ug mal/g	%	%	ppm	ppm	%	mg/100g
17. NaCl	18. NH ₃	19. SULFITE	20. Hg	21. TVB	22. DOMOIC ACID	23. TMA	24. VITAMIN A
%	mg/100g	ppm	ppm	mg N/100g	ug/ml	mgN/100g	IU/Qt
25. pH	26. BRIX	27. INSECT	28. RODENT	29. BENZOT'S	30. ASCORB'S	31. INDOLE	32. VITAMIN D
	%	%	%	ppm	ppm	ug/100g	IU/Qt
33. PKG INT	34. AFLATOXIN	35. Pb	36. Cd	37. Cu	38. Cr	39. Zn	40. A _w
	ppb	ppm	ppm	ppm	ppm	ppm	
41. Na	42. TSP	43. Ca	44. P	45. Mg	46. GLUTAMIC/MSG	47. ACETIC ACID	48. GLUCOSE
mg/Serving	mg/100g	mg/100g	mg/100g	mg/100g	%	mg/100g	%
47. FRUCTOSE	48. SUCROSE	49. LACTOSE	50. MALTOS	51. YEAST	52. MOLD	53. PSP	54. OTHER
%	%	%	%	%	%	%	%
56. FECAL/MPN	57. TOT COLIFORM/MPN	58. CL BOT	59. CL PERF	60. COAG STAPH	61. NON-COAG STAPH	62. LISTERIA	63. CAMPYLO
64. SPC	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC			
COLLECTOR'S NOTES: Case # 18 July 99				LABORATORY NOTES: Fe: 34.7			
EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)				ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE		CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY	
FOOD QUALITY (SAFETY SHELF-LIFE & CONSUMER ACCEPTANCE) INVOLVES THE ENTIRE CHAIN OF PRODUCTION PROCESSING & DISTRIBUTION OF FOOD*				REVIEWED BY:			
* SAMPLE WILL BE MAINTAINED FOR THIRTY (30) DAYS AFTER ANALYSIS AND THEN DISPOSED OF BY THE LABORATORY.				Supporting Documentation Task Force			
WHITE - DATA ENTRY				GOLDENROD - CONSUMER			

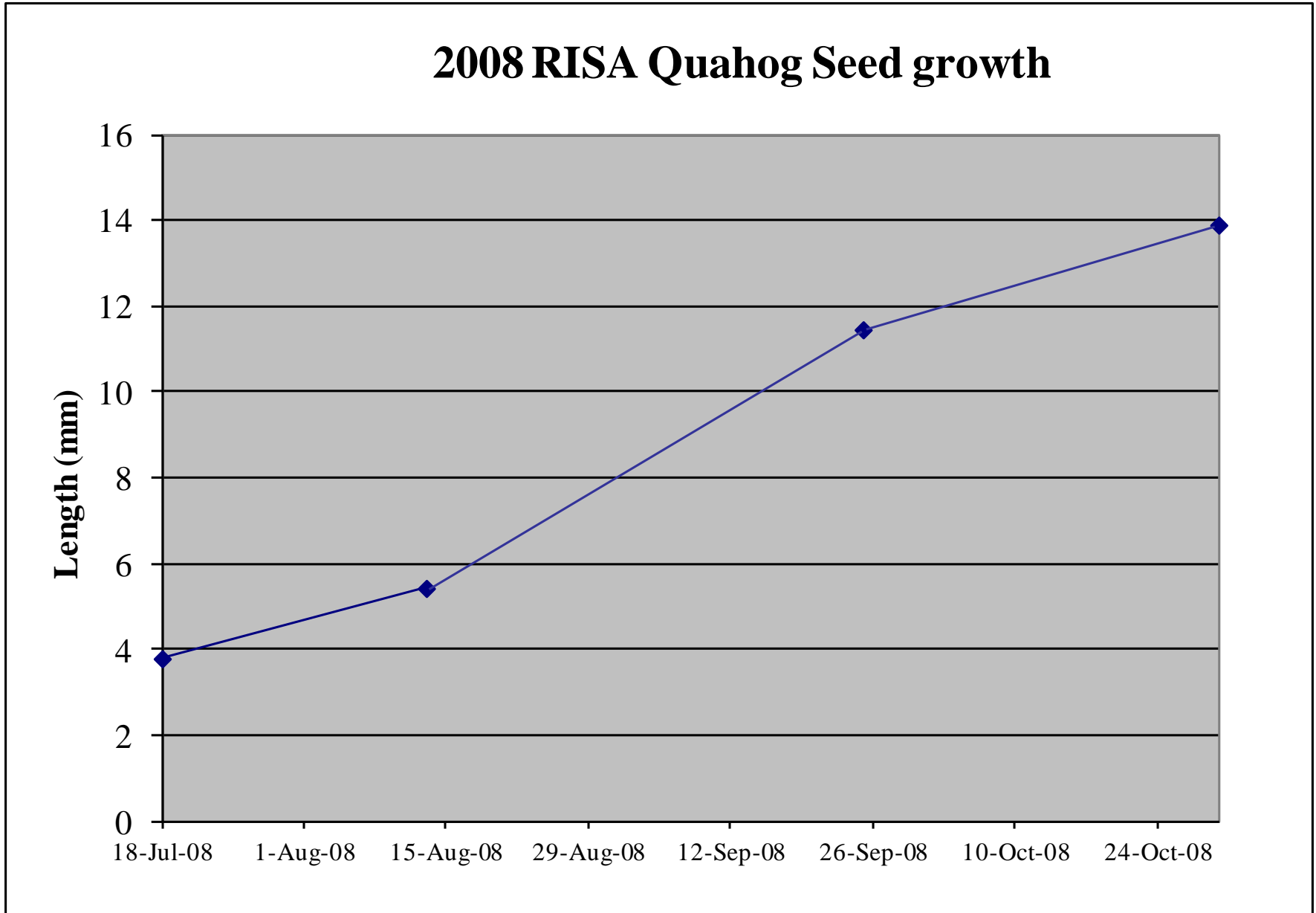


SAMPLE COLLECTION FORM

198249 13 AM 10:32

DATE OF COLLECTION 7/13/00 <i>ms</i>		DATE OF ANALYSIS 10/27/00	DATE OF REPORT 10/27/00	LAB SUP PCE			
CONDITION HOT FROZEN COLD OTHER	SEALED YES <input checked="" type="checkbox"/> NO <input type="checkbox"/>	ITEM OYSTERS	PRODUCT CODE/DOM	MOP	SAMPLE CODE 61		
EXP DATE	SIZE	BRAND NAME	TEMP ESTAB	TEMP REC °C	TEMP ANALYSIS		
DATE OF SHIP	FROM LOT OF	COLLECTED AT Moonstone Oyster	FOLLOW UP SAMPLE #1		REASON 01		
SHIPPER/PACKER & ADDRESS					SOURCE 02		
COMPLAINT*				ORIGINAL CONTAINER YES NO	ANALYSIS 13		
NAME & LOCATION OF STORE WHERE PURCHASED					DATE PURCHASED		
HOW STORED FROZEN COLD		AMBIENT	IMPORT PRODUCT YES NO	INTERVIEWED BY	DATE	TIME	AMOUNT REMAINING
CHAIN OF CUSTODY - DATE, TIME OF TRANSACTION OF COMPLAINT FROM FIELD TO LAB, ETC.							
SIGNED FROM <i>John Miller</i>	SIGNED TO <i>Mary Lou Silva</i>	DATE 7/13/00	TIME 10:35	HOW STORED <i>ambient</i>			
AGENT PRINT NAME 164 John Miller	PRINT NAME MARY LOU SILVA						
SIGNED FROM	SIGNED TO						
PRINT NAME	PRINT NAME						
SIGNED FROM	SIGNED TO						
PRINT NAME	PRINT NAME						
1. CEREAL %	2. MEAT PROT %	3. TOT H ₂ O %	4. SOY FL %	5. NFDM %	6. ADD H ₂ O %	7. TOT PROT %	8. TOT FAT %
9. SORBATES ppm	10. TBA ug mal/g	11. FFA %	12. WT/VOL %	13. NaNO ₂ ppm	14. EL'TROPHO ug/ml	15. TOTAL SOLIDS mgN/100g	16. HISTAMINE mg/100g
17. NaCl %	18. NH ₃ mg/100g	19. SULFITE ppm	20. Hg ppm	21. TVB mg N/100g	22. DOMOIC ACID ug/ml	23. TMA mgN/100g	24. VITAMIN A IU/Ct
25. pH	26. BRIX %	27. INSECT %	28. RODENT %	29. BENZOT'S ppm	30. ASCORB'S ppm	31. INDOLE ug/100g	32. VITAMIN D IU/Ct
33. PKG INT	34. AFLATOXIN ppb	35. Pb <0.54 ppm	36. Cd 0.19 ppm	37. Cu 60.9 ppm	38. Cr <0.48 ppm	39. Zn 711 ppm	40. A _w
41. Na mg/Serving	42. TSP mg/100g	43. Ca mg/100g	44. P mg/100g	45. Mg mg/100g	46. GLUTAMIC/MSG %	47. ACETIC ACID mg/100g	48. GLUCOSE %
47. FRUCTOSE %	48. SUCROSE %	49. LACTOSE %	50. MALTOSE %	51. YEAST %	52. MOLD %	53. PSP mg/100g	54. OTHER %
56. FECAL/MPN	57. TOT COLIFORM/MPN	58. CL BOT	59. CL PERF	60. COAG STAPH	61. NON-COAG STAPH	62. LISTERIA	63. CAMPYLO
64. SPC	65. SALMONELLA	66. B CEREUS	67. YERSINIA	ORGANOLEPTIC			
COLLECTOR'S NOTES: <i>Cage #238</i> <i>Rept. 99</i>				LABORATORY NOTES: <i>Fe: 38.3 ppm</i>			
EVALUATION BY LAB: 1. NOT AN FP OBLIGATION 2. FP VIOLATION - NO VIOLATION 3. FP ACTION INDICATED 4. INSUFFICIENT SAMPLE FOR ANALYSIS 5. INSUFFICIENT INFORMATION FOR ANALYSIS 6. IMPROPER CHAIN OF CUSTODY 7. LAB UNABLE TO PERFORM TESTING (REASON)				ADMINISTRATION FOLLOW-UP: 1. CHECK ESTABLISHMENT 2. ADDITIONAL SAMPLES 3. EMBARGO 4. REVIEW PROCESS 5. NO FURTHER ACTION 6. DISPOSE		CONCLUSION: 1. SATISFACTORY 2. QUESTIONABLE 3. VIOLATES STD 4. UNSATISFACTORY	
FOOD QUALITY (SAFETY SHELF-LIFE & CONSUMER ACCEPTANCE) INVOLVES THE ENTIRE CHAIN OF PRODUCTION PROCESSING & DISTRIBUTION OF FOOD*				REVIEWED BY:			
* SAMPLE WILL BE MAINTAINED FOR THIRTY (30) DAYS AFTER ANALYSIS AND THEN DISPOSED OF BY THE LABORATORY.							





Warwick Cove Upweller																		
Quahog Seed		30-Oct-08		Note: µg/kg = ppb														
Group (n=15)	avg Length (mm)	stdev	avg Live Weight (g)	stdev	avg Soft Tissue Wet Weight (g)	stdev			avg Hg/Soft Tissue Wet Weight (µg/kg)	stdev	avg Cr/Soft Tissue Wet Weight (µg/kg)	stdev	avg Fe/Soft Tissue Wet Weight* (µg/kg)	stdev	avg Ni/Soft Tissue Wet Weight (µg/kg)	stdev	avg Cu/Soft Tissue Wet Weight (µg/kg)	stdev
1	15.1	2.9	0.967	0.730	0.267	0.238			7.81		0.20		35.57		0.20		8.18	
2	12.6	1.6	0.545	0.202	0.139	0.056			9.41		0.28		34.00		0.22		11.80	
3	13.9	1.2	0.685	0.201	0.182	0.058			8.24		0.26		33.33		0.20		9.30	
Total	13.9	2.2	0.732	0.476	0.196	0.152			8.49	0.83	0.25	0.04	34.30	1.15	0.21	0.01	9.76	1.85
Group (n=15)					avg Soft Tissue Dry Weight (g)	stdev	avg % Dry Weight	stdev	avg Hg/Soft Tissue Dry Weight (µg/kg)	stdev	avg Cr/Soft Tissue Dry Weight (µg/kg)	stdev	avg Fe/Soft Tissue Dry Weight* (µg/kg)	stdev	avg Ni/Soft Tissue Dry Weight (µg/kg)	stdev	avg Cu/Soft Tissue Dry Weight (µg/kg)	stdev
1					0.041	0.041	14.8%	2.4%	52.75		1.38		240.70		1.37		55.33	
2					0.022	0.008	16.1%	1.5%	58.45		1.73		210.60		1.35		73.09	
3					0.027	0.008	15.2%	1.7%	54.22		1.70		219.20		1.31		61.16	
Total					0.030	0.025	15.4%	2.0%	55.14	2.96	1.60	0.19	223.50	15.50	1.34	0.03	63.19	9.05
Group (n=15)	avg Zn/Soft Tissue Wet Weight* (µg/kg)	stdev	avg As/Soft Tissue Wet Weight (µg/kg)	stdev	avg Se/Soft Tissue Wet Weight (µg/kg)	stdev	avg Sr/Soft Tissue Wet Weight (µg/kg)	stdev	avg Ag/Soft Tissue Wet Weight (µg/kg)	stdev	avg Cd/Soft Tissue Wet Weight (µg/kg)	stdev	avg Pb/Soft Tissue Wet Weight (µg/kg)	stdev				
1	55.10		1.77		0.87		15.28		0.03		0.06		0.31					
2	66.07		1.99		1.29		14.07		0.03		0.05		0.28					
3	55.07		1.55		0.53		11.94		0.03		0.11		0.26					
Total	58.75	6.34	1.77	0.22	0.89	0.38	13.76	1.69	0.03	0.00	0.07	0.03	0.28	0.03				
Group (n=15)	avg Zn/Soft Tissue Dry Weight* (µg/kg)	stdev	avg As/Soft Tissue Dry Weight (µg/kg)	stdev	avg Se/Soft Tissue Dry Weight (µg/kg)	stdev	avg Sr/Soft Tissue Dry Weight (µg/kg)	stdev	avg Ag/Soft Tissue Dry Weight (µg/kg)	stdev	avg Cd/Soft Tissue Dry Weight (µg/kg)	stdev	avg Pb/Soft Tissue Dry Weight (µg/kg)	stdev				
1	372.90		12.01		5.89		103.40		0.22		0.43		2.09					
2	409.30		12.32		7.97		87.14		0.16		0.29		1.76					
3	362.20		10.17		3.47		78.55		0.22		0.69		1.68					
Total	381.47	24.69	11.50	1.16	5.78	2.26	89.70	12.62	0.20	0.04	0.47	0.20	1.84	0.22				

Name of the New Method	MPN-Real-Time PCR Method for the Detection of <i>Vibrio Vulnificus</i> from Oysters
Name of the Method Developer	USFDA Gulf Coast Seafood Laboratory
Developer Contact Information	USFDA Gulf Coast Seafood Laboratory, 1 Iberville Drive, Dauphin Island, AL 36528 Jessica.jones@fda.hhs.gov

Checklist	Y/N	Submitter Comments
A. Need for the New Method		
Clearly define the need for which the method has been developed.	Y	<p>Currently, the most common NSSP method used to detect <i>Vibrio vulnificus</i> (Vv) in oysters is MPN-culture. The method is time consuming and laborious taking a minimum of four full days to produce a result. A quicker method uses Real-time PCR for detection, currently the only NSSP approved Real-time PCR utilizes Sybr green: a non-specific DNA binding molecule, which negates the ability to multiplex thus is not permissive of the use of an internal control to assure the reaction integrity. The ability to use an internal control adds a level of reliability the use of a non-specific binder like Sybr Green cannot. Additionally, the Sybr Green method is validated for use with the Smart Cycler by Cepheid which, as of December 2018, will no longer be supported by the manufacturer.</p> <p>The MPN Real-time PCR method for Vv detection in oysters will utilize the AB7500 Fast, the same instrument which the NSSP-approved MPN Real-time PCR methods for Vp utilizes. Further, this method uses a specific probe targeting the vvh gene of Vv and includes an internal control in a single assay. This assay is rapid and robust producing highly reliable results in 24-36 hours.</p>
What is the intended purpose of the method?	Y	Approved NSSP method for enumeration of Vv from oysters.

Is there an acknowledged need for this method in the NSSP?	Y	There is current methodology. This assay is quicker than the approved culture methods and more robust than the existing real-time PCR method.
What type of method? i.e. chemical, molecular, culture, etc.	Y	MPN enrichment with molecular confirmation.

B. Method Documentation		
1. Method documentation includes the following information:		
Method Title	Y	MPN-Real-Time PCR Method for the Detection of <i>Vibrio vulnificus</i> from Oysters
Method Scope	Y	This method is for the detection of <i>Vibrio vulnificus</i> from oysters using the AB7500 Fast real-time PCR platform.
References	Y	<p>Campbell, M.S., Wright, A.C., 2003. Real-time PCR analysis of <i>Vibrio vulnificus</i> from oysters. <i>Appl Environ Microbiol</i> 69, 7137-7144</p> <p>Jones, J.L., Kinsey, T.P., Johnson, L.W., Porso, R., Friedman, B., Curtis, M., Wesighan, P., Schuster, R., Bowers, J.C., 2016. Effects of Intertidal Harvest Practices on Levels of <i>Vibrio parahaemolyticus</i> and <i>Vibrio vulnificus</i> Bacteria in Oysters. <i>Appl Environ Microbiol</i> 82, 4517-4522.</p> <p>Kaysner, C., DePaola, A., 2004. <i>Vibrio</i>, Bacteriological Analytical Manual, 8th ed.</p> <p>Nordstrom, J.L., Vickery, M.C., Blackstone, G.M., Murray, S.L., DePaola, A., 2007. Development of a multiplex real-time PCR assay with an internal amplification control for the detection of total and pathogenic <i>Vibrio parahaemolyticus</i> bacteria in oysters. <i>Appl Environ Microbiol</i> 73, 5840-5847.</p>
Principle	Y	This method uses an MPN format for enumeration based on molecular (PCR) detection of the <i>vvh</i> gene specific to Vv.

Any proprietary aspects	Y	The AB7500 Fast is a proprietary real-time PCR platform developed by Applied Biosystems and sold through ThermoFisher Scientific. The optical plates and caps or film used are proprietary to the instrument.
Equipment required	Y	Equipment is listed in Appendix A.
Reagents required	Y	Media and reagents are listed in Appendix B.
Sample collection, preservation and storage requirements	Y	<p>Shellstock samples are bagged immediately upon collection and labeled with collector's name, the source of harvest, sampling stations, time, and date. Samples are placed in dry storage (ice chest or equivalent) maintained between 2°C and 10°C with ice or cold packs for transport. A layer of towels, bubblewrap, or another appropriate substance will separate shellfish from contact with ice or cold packs. If collected samples are frozen (such as IQF), direct contact with ice or cold packs is not permitted.</p> <p>Immediately upon arrival of sample(s) to the laboratory, date, time, and initials of receiver are documented. The temperature of three shellfish, each from a separate location within each shipping container, is measured by opening the shell enough to insert a temperature probe into the meat of the shellfish. If IQF samples are received, assure samples are frozen. Store at less than -15°C until ready to process. Temperatures are taken immediately after defrosting as described above. The shellfish is discarded after temperature is measured. Once temperature of the samples upon intake is established, the samples are placed under refrigeration for not longer than 36h after collection, unless processed immediately. Storage is documented. If processing IQF samples, samples are defrosted under refrigeration for no longer than 36h.</p>
Safety requirements	Y	Basic Personal Protection Equipment (PPE) is needed. A chain mail glove may

		be worn during shucking. Blending is done in a biosafety hood or the blender is placed in a splash shielded container or blender box. All biological waste is autoclaved and disposed of according to state regulations.
Clear and easy to follow step-by-step procedure	Y	Detailed procedure including sample preparation, MPN, PCR, and data analysis is included in Appendix C.
Quality control steps specific for this method	Y	Appropriately diluted process controls are used (Vv ATCC 33816 and Vp F11-3A). Appropriately diluted Internal Amplification Control (IAC) DNA is included in all PCRs. Manual review of amplification curves is conducted.

C. Validation Criteria		
1. Accuracy / Trueness	Y	Result: 110% Data: Table 1 Spike Range: -0.35 to 6.54 Log CFU/g
2. Measurement uncertainty	Y	Result: -0.57 to 0.044 log MPN/g Data: Table 1 Spike Range: -0.35 to 6.54 Log CFU/g
3. Precision characteristics (repeatability)	Y	Results: Variance ratio is <i>not significant</i> , based on least square regression. Calculated variability of the MPN method is 0.39, with a lower 95% CI of 0.32. The theoretical variability is 0.32. Data: Table 2, Figure 1 Spike Range: 0.38 to 5.54 Log CFU/g
4. Recovery	Y	Result: 110% Is the one way ANOVA to determine the consistency of recovery significant? No. Data: Table 2 Spike Range: 0.38 to 5.54 Log CFU/g
5. Specificity	Y	<i>V. alginolyticus</i> : $SI_{avg} = -1.28$, $p=0.42$ <i>V. cholerae</i> : $SI_{avg} = 1.26$, $p=0.09$ <i>V. fluvialis</i> : $SI_{avg} = -2.41$, $p=0.79$ <i>V. parahaemolyticus</i> : $SI_{avg} = 7.49$, $p=0.07$

		Data: Table 3 Range: 0.52 to 1.53 Log CFU/g
6. Working and Linear ranges	Y	Pearson's r: 0.97 Line equation: $\log(\text{MPN}) = 0.44 + 0.93 \times \log(\text{Plate Count})$ Is Pearson's r significant?: Yes Data: Table 4 and Figure 2 Range: -0.62 to 6.54 Log CFU/g
7. Limit of detection	Y	Result: 2.75 95% CI: 1.95, 3.88 Data: Table 4 and Figure 1 Range: -0.62 to 6.54 Log CFU/g
8. Limit of quantitation / Sensitivity	Y	Result: 0.3 MPN/g Data: Table 4 and Figure 1 Range: -0.62 to 6.54 Log CFU/g
9. Ruggedness	Y	Is there a significant difference between samples? Not under conditions tested. Data: Table 5 Range: 0.52 to 4.88 Log CFU/g
10. Matrix effects	Y	Effects of oyster matrix on the performance of the method was taken into consideration by using various sources of oysters for this study. Appendix D.
11. Comparability (if intended as a substitute for an established method accepted by the NSSP)	Y	No statistically significant difference between test and accepted methods. (p<0.05) Data: Table 6

D. Other Information		
1. Cost of the method	Y	Cost per sample for MPN: \$1.05 Cost per sample for PCR: \$20.55 Cost only includes reagents and consumables, infrastructure and personnel were not taken into account.
2. Special technical skills required to perform the method	Y	It is recommended that analysts have some formal training in molecular techniques or PCR, specifically.
3. Special equipment required and associated cost	Y	AB7500 FAST: \$34,060.00 AB7500 FAST annual maintenance contract: \$5,777.00
4. Abbreviations and acronyms defined	Y	Abbreviations and Acronyms are listed in appendix E.
5. Details of turn around times	Y	Results can be reported within 28h of

(time involved to complete the method)		sample receipt.
6. Provide brief overview of the quality systems used in the lab	Y	The laboratory adheres to the quality system standards of FDA/CFSAN, as well as those of the NSSP.

Submitters Signature	Date:
Submission of validation data and draft method to committee	Date:
Reviewing members:	
Accepted	Date:
Recommendations for further work	Date:

A. Validation Criteria

Data were generated using 20 separate lots of PHP oysters spiked with appropriate dilution(s) of a log phase culture of *Vibrio vulnificus*. Spike levels were determined by plate counts on TSA. Unless otherwise stated data was handled and analyzed as recommended in the SLV Documents for MPN Based Microbiological Methods on the ISSC website, with the exception of correcting for background using the blank sample data. The correction was not made because the levels in the blank samples were extremely low (near the LOD) and the it was more appropriate, from a statistical perspective, to not make the adjustment. For samples not detected, ½ the theoretical LOD was substituted for those values. For samples greater than the upper limit of the test, the values for the upper limit was used.

Table 1. Data used for determination of Accuracy/Trueness and Measurement Uncertainty.

Sample	Plate Count (Log CFU/g)	Sample Blank (Log MPN/g)	Spiked Sample (Log MPN/g)
1	-0.35	0.31	-0.25
2	4	ND*	3.33
3	1.19	ND	1.4
4	2.92	-0.45	3.17
5	1.38	ND	1.8
6	1.06	-0.52	2.36
7	2.74	-0.52	2.64
8	4.78	ND	4.96
9	4.84	ND	5.75
10	3	ND	3.38
11	6.54	ND	6.16
12	1.11	ND	1.63
13	6.08	0.36	5.36
14	4.88	ND	5.62
15	-0.19	ND	-0.15
16	2.57	ND	2.36
17	0.97	ND	1.92
18	1.53	ND	1.17
19	1.88	-0.45	2.16
20	0.52	-0.13	0.50

*ND=Not Detected

Table 2. Data used for determination of Precision and Recovery. Samples A and B are replicate analyses of the spiked homogenate.

Sample	Plate Count (Log CFU/g)	Sample Blank (Log MPN/g)	Spiked Sample A (Log MPN/g)	Spiked Sample B (Log MPN/g)
1	0.66	0.31	0.87	1.36
1	2.66	0.31	2.96	2.63
1	4.66	0.31	4.63	5.31
3	1.19	ND	1.63	1.17
3	3.19	ND	1.72	2.96
3	5.19	ND	3.96	5.17
5	0.38	ND	0.63	0.96
5	2.38	ND	2.63	2.17
5	4.38	ND	3.96	4.36
7	0.74	-0.52	1.32	1.31
7	2.74	-0.52	2.96	2.32
7	4.74	-0.52	4.97	5.35
9	0.84	ND	2.04	1.33
9	2.84	ND	3.66	3.38
9	4.84	ND	5.87	5.62
11	1.54	ND	1.63	1.63
11	3.54	ND	3.63	3.36
11	5.54	ND	5.34	5.62
13	1.08	0.36	1.36	1.96
13	3.08	0.36	2.96	3.16
13	5.08	0.36	5.18	4.97
15	0.81	ND	0.63	0.87
15	2.81	ND	2.63	3.96
15	4.81	ND	4.97	4.63
17	0.97	ND	1.97	1.87
17	2.97	ND	4.38	3.97
17	4.97	ND	5.87	5.887
19	0.88	-0.45	1.63	0.87
19	2.88	-0.45	3.36	3.36
19	4.88	-0.45	5.62	5.62

*ND=Not Detected

Figure 1. Plot of data from Table 2 by different concentrations (Low, Medium, and High). An alternative approach from ISSC recommendations to evaluating precision was used as a generalized least square regression with heterogenous variance structure was deemed a more appropriate test to estimate variance components for method error at different concentrations and then test whether or not method error varies significantly by concentration level. The output estimates of the variance components of the fit of two different models and then a comparison of those fits. One model has different parameters for method variation for each level (L, M, H) and the other constrains that variation to be the same across levels. The 1st model (null) estimates a common method error SD as 0.387 (same as the nested ANOVA). The 2nd model (full) estimates different method error SDs as 0.3217, 0.4688 and 0.3558 at levels L, M, and H respectively. Both models fit the same main effects (Levels nested within Samples) to remove that variation from what remains to determine method error estimates. A likelihood ratio test is used to compare the difference in the fit between the two models. The test statistic is the likelihood ratio between the two models and this is distributed as a Chi-square with 2 degrees of freedom (the difference in the number of parameters between the two models, 3 vs 1 variance parameters). The test statistic has a value of 1.58 and the p-value is 0.54 indicating no significant difference between the fits and hence no strong statistical evidence that method error varies across levels (L, M, H). The MSE for the residuals is 0.15. This corresponds to a SD of 0.39, which is only slightly higher than the theoretical method error SD (0.32), with a lower 95% confidence limit of 0.32.

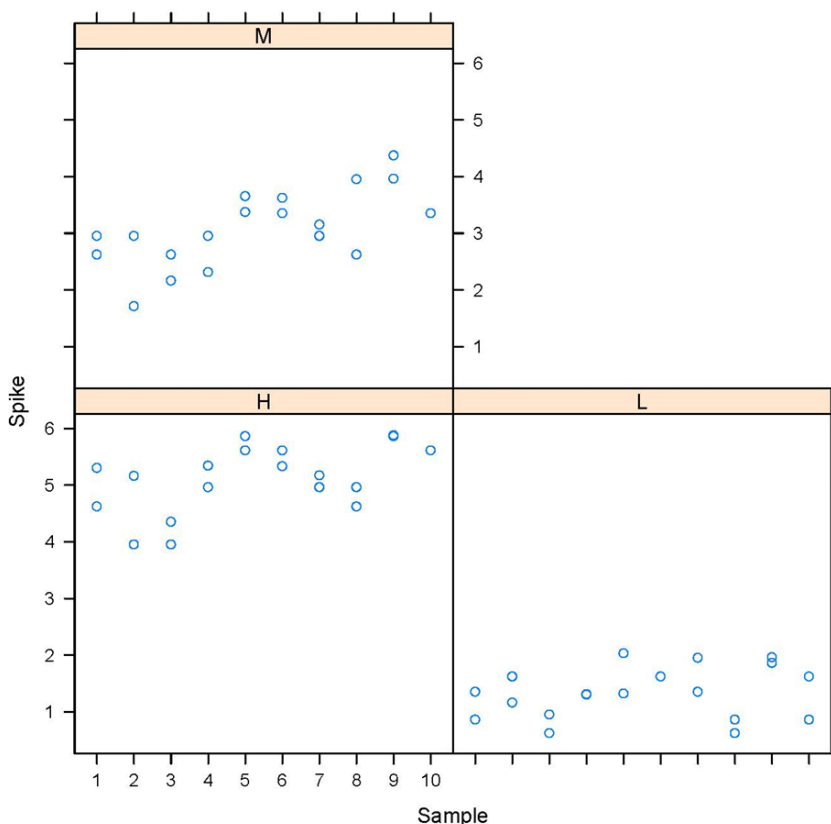


Table 3. Data used for determination of Specificity. Spike samples A-E are replicate analyses of the homogenate spiked only with Vv. Dual spike samples A-E are replicate analyses of the same homogenate spiked with Vv and the interfering organism.

Sample	Interfering Organism	Interfering Organism Plate Count (Log CFU/g)	<i>Vibrio vulnificus</i> Plate Count (Log CFU/g)	Sample Blank (Log MPN/g)	Spike Sample A (Log MPN/g)	Spike Sample B (Log MPN/g)	Spike Sample C (Log MPN/g)	Spike Sample D (Log MPN/g)	Spike Sample E (Log MPN/g)	Dual Spike Sample A (Log MPN/g)	Dual Spike Sample B (Log MPN/g)	Dual Spike Sample C (Log MPN/g)	Dual Spike Sample D (Log MPN/g)	Dual Spike Sample E (Log MPN/g)
6	<i>Vibrio parahaemolyticus</i>	5.49	1.06	-0.52	2.36	2.38	2.38	2.38	2.66	1.63	0.31	0.19	1.32	0.19
12	<i>Vibrio cholerae</i>	6.75	1.11	ND	1.63	1.63	1.63	1.63	1.96	1.36	0.96	1.36	1.63	1.63
18	<i>Vibrio fluvialis</i>	6.83	1.53	ND	1.17	1.36	1.63	1.36	1.86	1.96	1.96	1.63	-0.03	0.06
20	<i>Vibrio alginolyticus</i>	6.17	0.52	-0.13	0.45	0.96	0.31	0.45	0.45	0.96	0.44	0.43	-0.04	0.3

Table 4. Data used for determination of Working and Linear Ranges, Limit of Detection, and Limit of Quantitation/Sensitivity. Samples A and B are replicate analyses of the spiked homogenate. The LOQ is determined by the amount of inoculum used in the lowest dilution of the MPN, so long as the LOD is not statistically different than 1. As tested with a starting inoculum of 1g, the LOD of this method is 0.3 MPN/g

Sample	Plate Count (Log CFU/g)	Sample Blank (Log MPN/g)	Spike Sample A (Log MPN/g)	Spike Sample B (Log MPN/g)
1	-0.35	0.31	-0.45	-0.04
1	0.66	0.31	0.87	1.36
1	1.66	0.31	1.63	3.06
1	2.66	0.31	2.96	2.63
1	4.66	0.31	4.63	5.31
1	5.66	0.31	5.92	6.16
3	0.19	ND	0.86	0.63
3	1.19	ND	1.63	1.17
3	2.19	ND	2.36	2.87
3	3.19	ND	1.72	2.96
3	5.19	ND	3.96	5.17
3	6.19	ND	6.16	6.16
5	-0.62	ND	0	-0.45
5	0.38	ND	0.63	0.96
5	1.38	ND	1.63	1.96
5	2.38	ND	2.63	2.17
5	4.38	ND	3.96	4.36
5	5.38	ND	4.97	4.63
7	-0.25	-0.52	0.87	0.19
7	0.74	-0.52	1.32	1.31
7	1.74	-0.52	1.96	2.96
7	2.74	-0.52	2.96	2.32
7	4.74	-0.52	4.97	5.35
7	5.74	-0.52	5.92	6.16
9	-0.15	ND	0.31	0.17
9	0.84	ND	2.04	1.33
9	1.84	ND	1.87	2.04
9	2.84	ND	3.66	3.38
9	4.84	ND	5.87	5.62
9	5.84	ND	5.87	6.16
11	0.54	ND	0.36	0.63
11	1.54	ND	1.63	1.63
11	2.54	ND	2.36	2.87
11	3.54	ND	3.63	3.36

11	5.54	ND	5.34	5.62
11	6.54	ND	6.16	6.16
13	0.08	0.36	0.36	0.63
13	1.08	0.36	1.36	1.96
13	2.08	0.36	2.36	2.17
13	3.08	0.36	2.96	3.16
13	5.08	0.36	5.16	4.97
13	6.08	0.36	6.16	4.56
15	-0.19	ND	-0.45	0.16
15	0.81	ND	0.63	0.87
15	1.81	ND	1.45	0.54
15	2.81	ND	2.63	3.96
15	4.81	ND	4.97	4.63
15	5.81	ND	5.92	6.16
17	-0.03	ND	0.96	1.17
17	0.97	ND	1.96	1.87
17	1.97	ND	2.97	2.97
17	2.97	ND	4.38	3.97
17	4.97	ND	5.87	5.87
17	5.97	ND	6.16	6.16
19	-0.12	-0.45	-0.04	0.17
19	0.88	-0.45	1.63	0.87
19	1.88	-0.45	1.96	2.36
19	2.88	-0.45	3.36	3.36
19	4.88	-0.45	5.62	5.62
19	5.88	-0.45	6.16	6.16

Figure 2. Plot of data from Table 4 for determination of LOD/LOQ.

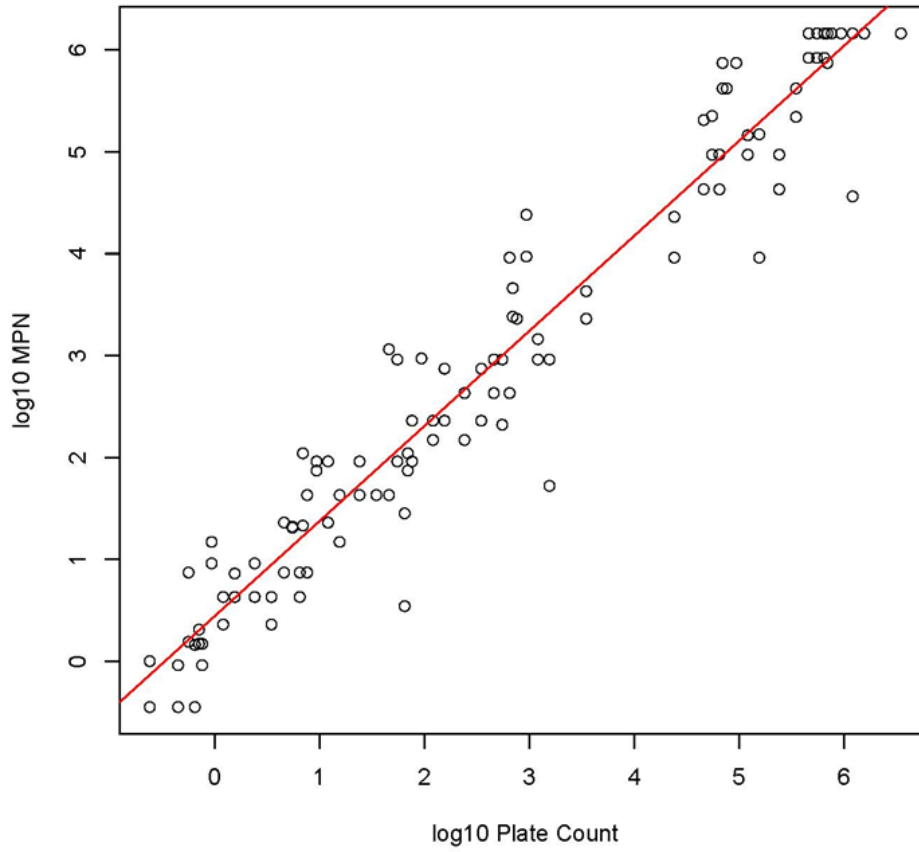


Table 5. Data used for determination of Ruggedness. Results reported as log MPN/g of *V. vulnificus* for each variation of the method SOP.

Sample	Plate Count (Log CFU/g)	Sample Blank (Log MPN/g)	Media/ Reagents		Oysters shucked and held prior to blending				RT 30 m Post-blending	MPN incubation			Boil prep time		Master Mix stored frozen				Master Mix thawed and re-frozen			Master Mix at RT 0.5-1.5h	
			Lot 1	Lot 2	4C 1h	4C 3h	RT 30m	RT 1h		35C >24h	RT 18-24h	39C 18-24h	5m	30m	5d	3d	2d	1d	4X	3X	2X		
2	4.00	0.52	3.33	5.04	3.38	4.04	3.66	4.04	4.04	5.04	4.04	3.66	4.04	4.04	3.33	3.06	3.33	3.33	3.33	3.33	3.33	3.33	2.54
4	2.92	-0.45	3.17	3.17	2.66	3.04	3.38	2.36	3.17	2.97	3.38	3.38	3.17	3.38	3.17	3.17	3.17	3.17	3.17	3.17	3.17	3.17	3.17
6	1.06	-0.52	2.36	2.36	0.06	-0.45	0.31	1.87	0.36	0.54	1.53	0.63	2.66	2.38	2.66	2.38	2.38	2.38	2.38	2.66	2.66	2.38	2.38
8	4.78	ND	4.96	4.63	4.38	4.17	4.38	4.63	4.63	4.59	4.38	4.63	4.63	4.96	4.63	4.63	4.63	4.63	4.63	4.63	4.63	4.63	4.63
10	3.00	ND	3.38	3.38	2.97	2.38	3.66	2.63	3.16	3.17	2.96	2.63	3.36	3.36	3.36	3.36	3.36	3.36	3.36	2.96	3.36	3.36	3.36
12	1.11	ND	1.63	1.63	1.63	0.87	1.86	1.36	1.63	1.3	1.17	1.16	1.63	1.63	1.96	1.63	1.63	1.63	1.63	1.63	1.63	1.63	1.63
14	4.88	ND	5.62	5.34	4.97	4.97	4.97	4.97	4.97	5.34	4.63	4.63	5.62	5.62	5.62	5.62	5.62	5.62	5.62	5.62	5.62	5.62	-0.52
16	2.57	ND	2.36	2.36	1.36	1.96	2.17	2.36	2.63	2.63	2.63	3.36	2.36	2.36	2.36	2.36	2.36	2.36	2.36	2.36	2.36	2.36	2.36
18	1.53	ND	1.17	1.36	1.63	1.36	1.86	1.36	1.63	1.63	1.63	2.17	1.32	1.63	1.96	1.63	1.63	1.96	1.96	1.96	1.96	1.96	1.96
20	0.52	-0.13	0.45	0.96	0.31	-0.04	0.96	0.19	-0.04	1.17	0.58	0.36	0.32	0.17	0.45	0.45	1.63	0.45	0.45	1.63	0.45	0.45	0.45

**PUBLIC HEALTH SERVICE
 U.S. FOOD AND DRUG ADMINISTRATION
 OFFICE OF FOOD SAFETY
 SHELLFISH AND AQUACULTURE POLICY BRANCH
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 TEL. 240- 402-2151/2055/4960 FAX 301-436-2601
 CFSANDSSLEOS@FDA.HHS.GOV**

SHELLFISH LABORATORY EVALUATION CHECKLIST

LABORATORY:

ADDRESS:

TELEPHONE:

FAX:

EMAIL:

DATE OF EVALUATION:

DATE OF REPORT:

LAST EVALUATION:

LABORATORY REPRESENTED BY:

TITLE:

LABORATORY EVALUATION OFFICER:

SHELLFISH SPECIALIST:

OTHER OFFICIALS PRESENT:

TITLE:

Items which do not conform are noted by: Conformity is noted by a “√”
 C- Critical K - Key O - Other NA- Not Applicable

Check the applicable analytical methods:

	MPN Real-time PCR method for <i>Vibrio vulnificus</i> detection in Oysters [PART III] SmartCycler II
	MPN Real-time PCR method for <i>Vibrio parahaemolyticus</i> detection in Oysters [PART III] SmartCycler II and AB 7500 Fast
	<u>MPN Real-time PCR method for <i>Vibrio parahaemolyticus</i> and <i>Vibrio vulnificus</i> detection in Oysters [Part III]</u>

PART I – Quality Assurance		
ITEM		
CODE	REF	
1.1 Quality Assurance (QA) Plan		
K	4, 6	1.1.1 Written Plan (Check <input checked="" type="checkbox"/> those items which apply).
		a. Organization of the Laboratory.
		b. Staff training requirements.
		c. Standard operating procedures (SOPs).
		d. Internal quality control measures for equipment, their calibration maintenance, repair, performance and rejection criteria established.
		e. Laboratory safety.
		f. Internal performance assessment.
		g. External performance assessment.
C	4	1.1.2 The QA plan is implemented.
K	6	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify the program(s): _____
1.2 Educational/Experience Requirements		
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor must have at least a bachelor's degree in microbiology, biology or equivalent discipline with at least two years of laboratory experience.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analysts meet the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree in microbiology, biology or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial laboratories, the analysts must have at least a high school diploma and at least three months of experience in laboratory sciences.
1.3 Work Area		
O	4, 6	1.3.1 Adequate for workload and storage.
K	6	1.3.2 Clean, well lighted.
K	6	1.3.3 Adequate temperature control.
O	6	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	6	1.3.5 Microbiological quality of the air contains fewer than 15 colonies/plate for a 15 minute exposure determined monthly. The results are recorded and records maintained.
1.4 Laboratory Equipment		
K	5	1.4.1 To determine the pH of prepared media and reagents, the pH meter has a standard accuracy of 0.1 pH units.
K	9	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	6	1.4.3 The effect of temperature on the pH is compensated for by an internal/external ATC probe or by manual adjustment (<i>Circle the appropriate type of adjustment</i>).
K	4	1.4.4 The pH meter is calibrated daily or with each use as per product literature. Results are recorded and records maintained.
K	6	1.4.5 A minimum of two standard buffer solutions are used to calibrate the pH meter. The first is near the electrode isopotential point (pH 7). The second is near the expected sample pH (i.e. pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	4	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt procedure or through determination of the slope (<i>Circle the method used</i>).
K	5	1.4.7 The balances used provide a sensitivity of at least 0.1 g at the weights of use.

K	6		1.4.8 Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	6		1.4.9 Refrigerator temperatures are monitored at least once daily on workdays. Results are recorded and records maintained.
K	1		1.4.10 Refrigerator temperatures are maintained between 0 and 4 °C, except for reagent refrigerators which are maintained between 2 and 8 °C.
C	7		1.4.11 Freezer temperature is maintained at -15 °C or below.
O	7		1.4.12 Freezer temperature is monitored at least once daily on workdays. Results are recorded and records maintained.
C	5		1.4.13 The temperature of the incubator is maintained at 35 +/- 2.0 °C.
K	6		1.4.14 Thermometers used in the air incubators are graduated at no greater than 0.5 °C increments.
K	5		1.4.15 Working thermometers are located on top and bottom shelves of use in the air incubator or appropriately placed based on the results of spatial temperature checks.
K	4, 6		1.4.16 Air incubator temperatures are taken twice daily on workdays. Results are recorded and records maintained.
C	3		1.4.17 All working thermometers are appropriately immersed.
C	2, 20		1.4.18 Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	6, 20		1.4.19 A standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0 and 35. These calibration records are maintained.
K	3, 5		1.4.20 Standard thermometers are checked annually for accuracy by ice point determination. Results are recorded and maintained. Date of most recent determination: _____
C	2, 20		1.4.21 Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of ≤ 0.05 °C are used as the laboratory standards thermometer (Circle the thermometer type used).
K	3, 8		1.4.22 All working thermometers are checked annually against the standards thermometer at temperature(s) of use. Results are recorded and records maintained.
O	6		1.4.23 Appropriate pipet aids are available and used to inoculate samples.
K	2		1.4.24 Micropipettors are calibrated annually at appropriate volumes used and checked for accuracy quarterly. Results are recorded and records maintained.
1.5 Labware and Glassware Washing			
K	5		1.5.1 Utensils, containers, glassware and plasticware are clean borosilicate glass, stainless steel or other noncorroding material.
K	5		1.5.2 Culture tubes are new and of a suitable size to accommodate the volume for nutritive ingredients and sample.
K	5		1.5.3 Dilution bottles and tubes are made of borosilicate glass or plastic and closed with secure caps or screw caps with nontoxic liners.
K	5		1.5.4 Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
K	5		1.5.5 In washing reusable pipets, glassware and labware, a succession of at least three fresh water rinses plus a final rinse of deionized water is used to thoroughly rinse off all detergent.
C	2		1.5.6 An alkaline or acidic detergent is used for washing glassware/labware.
C	6		1.5.7 With each load of labware/glassware washed, the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali as appropriate) with aqueous 0.04% bromothymol blue (BTB) solution. Results are recorded and records maintained.
1.6 Sterilization and Decontamination			

K	5		1.6.1 The autoclave is of sufficient size to accommodate the workload.
K	4		1.6.2 Routine autoclave maintenance is performed and the records maintained.
C	6, 20		1.6.3 The autoclave provides a sterilizing temperature of 121 ± 2 °C as determined for each load using a calibrated maximum registering thermometer. As an alternative, an appropriate temperature monitoring device is used in place of the maximum registering thermometer when these are unavailable due to the ban on mercury.
K	6		1.6.4 An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121 °C. Calibration at 100 °C, the steam point is also recommended but not required.
K	10		1.6.5 The autoclave standards thermometer is checked every five years for accuracy at either 121 °C or at 100 °C, the steam point if the thermometer has been previously calibrated at this temperature. Date of most recent determination: _____
K	1		1.6.6 Working autoclave thermometers are checked against the autoclave standards thermometer at 121 °C yearly. Date of last check: _____
K	6		1.6.7 Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	6		1.6.8 Heat sensitive tape is used with each autoclave batch.
K	6		1.6.9 Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings (<i>Circle the appropriate type or types</i>).
K	6		1.6.10 For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180 °C.
K	5		1.6.11 A thermometer capable of determining temperatures accurately in the range of 160 to 180 °C is used to monitor the operation of the hot air sterilizing oven.
K	8		1.6.12 Records of temperature and exposure times are maintained for the operation of the hot-air sterilizing oven.
K	6		1.6.13 Spore strips/suspensions appropriate for use in dry heat are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Results are recorded and records maintained.
K	5		1.6.14 Reusable pipets are stored and sterilized in aluminum or stainless steel containers.
K	5		1.6.15 Reusable pipets (in canisters) are sterilized in a hot-air oven at 170 °C for 2 hours.
C	2		1.6.16 The sterility of reusable pipets is determined with each load sterilized. Results are recorded and records maintained.
C	2		1.6.17 The sterility of autoclave sterilized disposable pipet tips and microcentrifuge tubes is determined with each load sterilized. Results are recorded and records maintained. If presterilized pipet tips and microcentrifuge tubes are purchased certificate should be maintained and sterility confirmed as in 1.6.18.
C	2		1.6.18 The sterility of presterilized disposable pipets, pipet tips and microcentrifuge tubes is determined with each lot received. Results are recorded and records maintained.
K	8		1.6.19 Spent broth cultures and agar plates are properly decontaminated before disposal.
			1.7 Media Preparation
K	13, 14		1.7.1 Alkaline peptone water (APW) is prepared from the individual components and pH adjusted appropriately.
K	6		1.7.2 Media components are properly stored in a cool dry place.
O	6		1.7.3 Media components are labeled with the analyst's initials, date of receipt and date opened.
O	6		1.7.4 Dehydrated media are labeled with date of receipt and date opened.
C	6		1.7.5 Caked or expired media or media components are discarded.

C	6		1.7.6 Reagent water for media and diluent preparation is analyzed for residual chlorine monthly and is at a non-detectable level (≤ 0.1 ppm). Results are recorded and records maintained
K	6		1.7.7 Reagent water for media and diluent preparation contains <100 CFU/mL as determined monthly using the heterotropic plate count method. Results are recorded and records maintained.
K	5		1.7.8 The volume and concentration of media in the tube is suitable for the amount of sample inoculated.
C	6		1.7.9 Media broths are not in the autoclave for more than 60 minutes.
C	1		1.7.10 Media and diluent sterility is determined for each load sterilized. Results are recorded and records maintained.
C	1		1.7.11 Media productivity is determined using media-appropriate positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is made from its individual components.
C	6		1.7.12 The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer requirements and/or method tolerance. Results are recorded and records are maintained.
			1.8 Storage of Prepared Culture Media
K	5		1.8.1 Prepared culture media are stored in a cool, clean, dry place where excessive evaporation and the danger of contamination is minimized.
K	8		1.8.2 Stored media are labeled with the storage expiration date or sterilization date.
K	5		1.8.3 Storage of prepared culture media at room temperature does not exceed 7 days.
K	2		1.8.4 Storage under refrigeration of prepared broth media with loose fitting closures does not exceed 1 month.
K	6		1.8.5 Storage under refrigeration of prepared culture media with screw- cap closures does not exceed 3 months.
K	11		1.8.6 All prepared broth media stored under refrigeration is warmed to room temperature prior to use, without exceeding incubation temperature.
PART II –Samples			
			2.1 Sample Collection, Transportation and Receipt
C	2, 6		2.1.1 A representative sample is collected and a chain of custody documenting the history of the sample(s) from collection to final disposal has been established.
K	5		2.1.2 Shellfish samples as received are collected in clean, waterproof, puncture resistant containers loosely sealed or are rejected for regulatory analysis.
K	5		2.1.3 Shellfish samples as received are labeled with the collector's (or if PHP, company/processor and collector's) name, the source, the time and date of collection or are rejected for regulatory analysis.
C	5		2.1.4 Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 2 and 10 °C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1		2.1.5 Analysis of the samples is initiated as soon as possible after collection, but not to exceed 36 h. If processing IQF samples, samples are defrosted under refrigeration for no longer than 36 h once removed from the freezer.
			2.2 Preparation of Samples for Analysis
K	2, 6		2.2.1 Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes.
O	2		2.2.2 Blades of shucking knives are not corroded.
K	5		2.2.3 The hands of the analyst are thoroughly washed with soap and water or new gloves are donned, immediately prior to cleaning the shells of debris.
O	2		2.2.4 The faucet used for rinsing the shellfish does not contain an aerator.
K	5		2.2.5 Shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
K	5		2.2.6 Samples are allowed to drain in a clean container or on clean towels prior to opening
K	5, 15		2.2.7 Immediately prior to shucking, the hands or gloved hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol. The gloves if worn are latex, nitrile and/or stainless steel mesh to protect analyst's hands from injury.
C	5		2.2.8 Shellfish are not shucked through the hinge.

C	5		2.2.9 The contents of the sample (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
C	5		2.2.10 A representative sample of at least 12 shellfish is used for analysis
C	2, 5		2.2.11 A quantity of meat and liquor is sufficient to cover the blender blades or additional oysters are used in order to ensure sample homogeneity.
K	2, 13		2.2.12 The sample can be processed directly or a 1:1 dilution of shellfish:diluent made. If a dilution is made, the sample is weighed to the nearest 0.1 g and an equal amount, by weight, of diluent is added.
K	13		2.2.13 Sterile phosphate buffered saline (pH 7.4) is used as the sample diluent.
C	5		2.2.14 Samples are blended for 60 to 120 seconds until homogenous.
PART III- PCR method for <i>Vibrio vulnificus</i> and <i>Vibrio parahaemolyticus</i> detection in Oysters			
3.1 APW Enrichment			
K	5		3.1.1 Sterile phosphate buffered saline (PBS) is used as the sample diluent.
C	5, 15		3.1.2 The 1:10 dilution is prepared gravimetrically with PBS. All successive dilutions are prepared volumetrically. For example, if an initial 1:1 dilution of the sample was used for blending, the 1:10 dilution is prepared by adding 20 g of sample homogenate to 80 ml of PBS. If the homogenate was not diluted, the 1:10 dilution is prepared by adding 10 g of sample homogenate to 90 ml of PBS.
C	17		3.1.3 Appropriate sample dilutions are inoculated into APW. Specify dilution(s) used _____ Specify number of tubes per dilution _____
C	2, 15 21		3.1.4 For <i>V. parahaemolyticus</i> analysis, a tdh+, trh+ <i>V. parahaemolyticus</i> culture diluted to 10^3 per ml is used as a positive process control. A non <i>V. parahaemolyticus</i> culture is used as a negative process control. For <i>V. vulnificus</i> analysis, a <i>V. vulnificus</i> culture diluted to 10^3 per ml is used as a positive process control. A non <i>V. vulnificus</i> culture is used as a negative process control. <u>An uninoculated APW blank will serve as the uninoculated control.</u> The process control cultures accompany the samples throughout incubation, isolation, and confirmation. Records are maintained.
C	13		3.1.5 Inoculated APW enrichment tubes are incubated at 35 +/- 2 °C.
C	13		3.1.6 Tubes are read after 18 – 24 hours of incubation. Clear tubes are negative. Turbid tubes are positive and shall be further processed.
3.2 PCR Reagents			
C	14, 15		3.2.1 Lyophilized primers and probes are stored according to manufacturer's instructions.
K	14, 15		3.2.2 Fluorescent probes are stored in light occluding tubes or containers.
C	14, 15, 18, 1921		3.2.3 The PCR forward and reverse primers and probes are appropriate for the platform target. <u>For Total and Pathogenic Vp Real-time PCR Method</u> <u>Trh 627F: 5' ATA CCT TTT CCT TCT CCW GGT TC 3'</u> tdh_269-20:—6FAM-5'-TGACATCCTACATGACTGTG-3' MGBNFQ <u>Trh 731b R: 5' TTG TCC AGT AGT CAT CAA CGA TTG 3'</u> trh_133-23: NED/TET-5'-AGAAATACAACAATCAAACTGA-3' MGBNFQ <u>Trh Glov R: 5' TTG TCC AAT AGT CCT CCA CAA TTG 3'</u> trh_1043: JOE/TEXAS-RED-5'-CGCTCGCGTTCACGAAACCGT-3' BHQ2 <u>WA IC F: 5' GGC GAA GCG AAT CTG GAA A 3'</u> IAC_109: CY5-5'-TCTCATGCGTCTCCCTGGTGAATGTG-3' BHQ2 <u>WA IC R: 5' GGT GTA GTT GTG CGT GTA ATA TGA GA 3'</u> trh_20F: 5'-TTGCTTTCAGTTTGCTATTGGCT-3'

		<p><u>Orf8 F: 5' TCA CCT GAG GAC GCA GTT ACG 3'</u>trh_292R: 5'-TGTTTACCGTCATATAGGCGCTT 3'</p> <p><u>Orf8 R: 5' TTC AAT TGT AGA ACC GCC AGC TA 3'</u>tdh_89F: 5'-TCCCTTTTCTGCCCCC 3'</p> <p><u>Tlh-F: 5' CCG CTG ACA ATC GCT TCT C 3'</u>tdh_321R: 5'-CGCTGCCATTGTATAGTCTTTATC 3'</p> <p><u>Tlh-R: 5' TTT GAT CTG GCT GCA TTG CT 3'</u>thh_884F: 5'-ACTCAACACAAGAAGAGATCGACAA 3'</p> <p><u>TDH-F: 5' TAT CCA TGT TGG CTG CAT TC 3'</u>thh_1091R: 5'-GATGAGCGGTTGATGTCCAAA 3'</p> <p><u>TDH-R: 5' CGA ACA ACA AAC AAT ATC TCA TCA GA 3'</u>IAC_46F: 5'-GACATCGATATGGGTGCCG 3'</p> <p><u>Trh Probe: 6FAM 5' TAT TTG TYG TTA GAA ATA CAA CAA T 3'</u></p> <p><u>MGBNFQ IAC_186R: 5' CGAGACGATGCAGCCATTC 3'</u></p> <p><u>WA IC Probe :VIC 5' CGT AAG ACA ATC TGA TAG TAG T 3' MGBNFQ Orf8 Probe: NED 5' TCC TGC TGT ACT TTT AG 3' MGBNFQ</u></p> <p><u>Tlh Probe: 6FAM 5' ACC ACA CGA TCT GGA GCA ACG ACG MGBNFQ TDH Probe 3' VIC TGT CCC TTT TCC TGC CCC CGG 5' MGBNFQ</u></p> <p>For Vv Real-time PCR Method</p> <p><u>vvha-F: 5' GAT CGT TGT TTG ACC GTA AAC G 3'</u></p> <p><u>vvha-R 5' TGC TAA GTT CGC ACC ACA CTG T 3'</u></p> <p><u>vvha Probe: NED 3' CAA AAC GCT CAC AGT CG 5' MGB probe</u></p> <p><u>vvhF 5' TGTTTATGGTGAGAACGGTGACA 3'</u></p> <p><u>vvhR 5' TTCTTTATCTAGGCCCAA ACTTG 3'</u></p>
C	14, 18	3.2.4 Lyophilized forward and reverse primers, and probes, are hydrated with TE buffer to produce a 0.1 mM stock solution.
C	<u>14,1821</u>	3.2.5 <u>Storage of thawed working stocks of primers and probes are stored between 2-8°C, not to exceed 2 weeks. Using molecular grade, nuclease free water, primer and probe stock solutions are diluted to produce a 0.01 mM working solution.</u>
C	<u>14,1821</u>	3.2.6 <u>Storage of aliquoted working stocks of Reconstituted primers and probes are stored in a -20 °C manual defrost freezer does not exceed 1 year, for up to 5 freeze-thaw cycles, not to exceed two years.</u>
C	<u>21,22</u>	3.2.7 <u>Taqman Environmental Mastermix 2.0 is stored in -20°C manual defrost freezer until first use. Platinum Taq DNA is stored in -20 °C manual defrost freezer until first use. After first use, can-beit is stored between 2-8 °C.</u>
C	<u>21,22</u>	3.2.8 <u>Internal control (IC) is stored in -20°C manual defrost freezer until first use. PCR reagents (dNTPs, buffer, MgCl2, fluorescent dyes) are stored in -20 °C manual defrost freezer until first use. After first use, they can-be are stored between 2-8 °C.</u>
3.3 DNA Extraction		
C	14, 18	3.3.1 All microcentrifuge tubes and pipet tips are sterile.
C	14, 18	3.3.2 Pipet tips have aerosol barriers.
K	14, 18	3.3.3 Latex or nitrile gloves are worn throughout the extraction and PCR preparation process.
K	14, 18	3.3.4 All work surfaces, centrifuge racks and equipment used in PCR analysis are disinfected immediately prior to DNA extraction, Master Mix preparation and PCR analysis.
C	14, 18	3.3.5 Aseptic technique is observed throughout the extraction and PCR analysis.
C	<u>14, 18, 21</u>	3.3.6 <u>Two-hundred (200) µL One thousand (1000) µL aliquots from each positive APW enrichment tube, including the process controls, are extracted.</u>
C	<u>14,1821</u>	3.3.7 <u>For each run a specified amount of internal control (IC) is prepared such that each extracted well contains internal control DNA. Positive APW aliquots are placed in sterile microcentrifuge tubes and heated at 95-100 °C for 10 minutes.</u>
K	<u>14,1821</u>	3.3.8 <u>Extracts are refrigerated between 2-8°C and analyzed within 24 hrs. Frozen extracts are analyzed within 1 month of frozen storage. A set of positive and negative process controls are included with each batch of samples in a heating block/boiling bath.</u>

C	<u>14, 18, 21</u>		3.3.9 <u>A tlh+ trh+ tdh+ V. parahaemolyticus (WA4647 or equivalent), a tlh+ tdh+ Orf8+ V. parahaemolyticus (BAA-240 or equivalent), and vvha+ V. vulnificus (ATCC 27562 or equivalent) cultures are extracted and combined to serve as the positive PCR (amplification) control. After boiling, tubes are chilled in ice or immediately frozen in a manual defrost freezer for future analysis. Boil preps may be refrigerated not to exceed 72 hours.</u>
K	<u>14, 18</u>		3.3.10 Frozen extracts are analyzed within 6 months of frozen storage.
3.4 Preparation of the Master Mix for PCR			
C	14, 16, 18		3.4.1 Nuclease-free microcentrifuge tubes and pipette tips, with filters, are used in Master Mix preparation.
C	<u>14, 16, 18, 21</u>		3.4.2 For each reaction, add the specified amount of water, buffer, MgCl₂, dNTPs, specific primers, nuclease probes, <i>Taq</i>, and internal control DNA is added.
K	14, 21 16, 18		3.4.3 The Master Mix is gently vortexed to mix constituents and then briefly spun <u>immediately prior to dispensing aliquots to reaction tubes or plates.</u>
C	14, 16, 18, <u>21</u>		3.4.4 <u>Eighteen (18) μL</u> Twenty-three (23) μL of Master Mix is used for each PCR reaction.
C	14, 16, 18		3.4.5 Master Mix must be used on the day of preparation or stored at –20 °C until time of use.
3.5 PCR			
C	<u>14, 19</u>		3.5.1 If previously frozen, the DNA extracts are completely thawed at temperatures no warmer than room temperature. Immediately prior to use, DNA extracts are centrifuged at >5,000 x g for 2 minutes to remove particulate matter and cell debris.
C	14, 19, <u>21</u>		3.5.2-1 Two (2) μL of DNA template is added to each reaction tube or plate well containing <u>23-18</u> μL of Master Mix for a total PCR reaction volume of <u>25-20</u> μL.
C	<u>14, 19, 21</u>		<u>3.5.2 Two (2) μL of extracted blank APW from the uninoculated process control is added to a reaction tube or plate well containing 18μL of Master Mix.</u>
K	14, 19, <u>21</u>		3.5.3 Two (2) μL of molecular grade, nuclease free water is added to a reaction tube or plate well containing 23-18 μL of Master Mix for each batch of Master Mix prepared as a no template control.
C	14, 19, <u>21</u>		3.5.4 Two (2) μL of DNA template extracted from the negative process control culture is added to a reaction tube or plate well containing <u>23-18</u> μL of Master Mix.
C	14, 19, <u>21</u>		3.5.5 Two (2) μL of DNA template extracted from the positive process control culture is added to a reaction tube or plate well containing <u>23-18</u> μL of Master Mix.
O	14, 19, <u>21</u>		3.5.6 Two (2) μL of DNA template extracted from the positive control culture (prepared separately from the positive process control) is added to a reaction tube or plate well containing 23-18 μL of Master Mix as the positive PCR (<u>amplification</u>) control.
K	14, 19, <u>21</u>		3.5.7 Immediately prior to loading the reaction tubes or plates into the instrument they are centrifuged for 3-5 <u>30</u> seconds to ensure that all reagents and the DNA template are in the bottom of the tube to optimize the PCR amplification process.
C	16		3.5.8 After centrifugation, tubes or plates are inserted into the instrument.
3.6 PCR Amplification			
C	14, 19		3.6.1 The appropriate instrument platform is used for the protocol.
K	16		3.6.2 Manufacturer’s instructions are followed in operating the instrument.
C	14, 19		3.6.3 The PCR cycle parameters used are appropriate for the protocol.
K	14, 19		3.6.4 Optical calibrations for the dyes being used are current, per the instrument manufacturer’s recommendations.
C	14, 19		3.6.5 The analysis settings are adjusted as specified in the protocol.
3.7 Computation of Results			
K	14, 19		3.7.1 All runs in which the positive control generates a Ct value for the target(s) of interest and the negative control reaction generates no Ct value for the target(s), but a Ct value for the internal control are considered valid.
C	2		3.7.2 Data is quality checked by the analyst.
C	14, 19		3.7.3 All reactions in a valid run which generate a Ct value for the target(s) of interest with a sigmoidal amplification curve are considered to be positive.

C	16		3.7.4 Any sample which does not demonstrate a sigmoidal amplification curve may have a reported positive/negative determination that is discrepant from the instrument if appropriately justified using the raw fluorescent data.
K	16		3.7.5 All reactions in a valid run which do not generate a Ct value for the target(s) of interest, but do generate a Ct value for the internal control are considered negative.
C	16		3.7.6 Any reaction in which no Ct value is generated for the target(s) of interest or the internal control is considered invalid and should be re-tested.
C	13		3.7.7 Upon determination of positive reactions, refer to the original positive dilutions of APW and record MPN values as derived from the calculator in Appendix 2 of the FDA Bacteriological Analytical Manual (BAM).
K	13		3.7.8 For APW enrichment, results are reported as MPN/g of sample.

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LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
MICROBIOLOGICAL COMPONENT: (Part I-III)	
A. Results	
Total # of Critical (C) Nonconformities in Parts I-III	
Total # of Key (K) Nonconformities in Parts I-III	
Total # of Critical, Key and Other (O)	
Nonconformities in Parts I-III	
B. Criteria for Determining Laboratory Status of the Microbiological Component:	
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <p style="margin-left: 40px;">a. The total # of Critical nonconformities is ≥ 4 or _____</p> <p style="margin-left: 40px;">b. The total # of Key nonconformities is ≥ 13 or _____</p> <p style="margin-left: 40px;">c. The total # of Critical, Key and Other is ≥ 18 _____</p> <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1</p>	
C. Laboratory Status (circle appropriate)	
Does Not Conform	Provisionally Conforms
Conforms	
Acknowledgment by Laboratory Director/Supervisor:	
All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____.	
Laboratory Signature: _____	Date: _____

1. Purpose/Principle

The purpose of this test is to rapidly quantify *Vibrio parahaemolyticus* (*Vp*) and *Vibrio vulnificus* (*Vv*) from oysters using a high throughput MPN based real-time PCR protocol. Culture based assays for the enumeration of *Vp* and *Vv* require a minimum of four days and require the Kanagawa test (media based) to detect pathogenicity. This assay was designed to provide quantitative results for total *Vibrio parahaemolyticus* (*Vp tlh+*), known pathogenic markers of *Vibrio parahaemolyticus* (*Vp tdh+* and *Vp trh+*), as well as a strain of potential pandemic *Vibrio* (*Vp ORF8+*). Additionally, the assay provides quantitative results for total *Vibrio vulnificus* (*vvhA*) and utilizes an exogenous internal control (WA IC).

This test utilizes Taqman® probe real-time polymerase chain reactions to amplify 4 target genes from the *Vibrio parahaemolyticus* (*Vp*) genome as well as 1 target from the *Vibrio vulnificus* (*Vv*) genome.

Vp

- Thermolabile hemolysin, *tlh* gene
- Thermostable direct hemolysin, *tdh* gene
- Thermostable direct related hemolysin, *trh* gene
- Filamentous phage (f237) Orf8, gene

Vv

- Cytolysin-hemolysin, *vvhA* gene

2. Scope

Rapid and early detection of these pathogens will help the shellfish industry market oysters for consumption that are within regulatory limits for these pathogens, and ensure public health safety.

3. Reagents / Media

- Master Mix: TaqMan™ Environmental Master Mix 2.0; Thermo Fisher Cat. #4396838
- Molecular PCR grade water; Thermo Fisher Cat. #SH3053802 or equivalent
- TE buffer; Thermo Fisher Cat. #BP2473500 or equivalent
- Primers (See appendix A for sequences)
- Probes (See appendix A for sequences)
- Internal Control Plasmid
- MagNAPure 96 DNA and Viral NA Small Volume kit; Roche, Cat. # 06543588001
- Alkaline Peptone Water (APW); Prepared In-house
- Phosphate Buffer Saline (PBS); Prepared In-house

Record receipt of all PCR mastermix components in the Reagent Receipt Log (P:\EHSPHL\PHL\MICRO\FOODLAB\Vibrio\Master Mix Documents\Updated Worksheets\Reagent Receipt Log). All reagents will be tracked by its lot number. The intent of the reagent receipt logbook is to allow for complete traceability. Additionally, lot numbers are recorded upon use on Master Mix Worksheet.

Prepare Primer and Probe mixes according to the Master Mix Worksheet (P:\EHSPHL\PHL\MICRO\FOODLAB\Vibrio\Master Mix Documents\Mastermix Template). Lyophilized primers are diluted to 100 μ M stock concentrations with TE Buffer and stored at -15°C (or below) until use. Store in low light transmitting tubes. Original stock solutions are good for 5 years unless otherwise stated by manufacturers. Working concentrations of primers and probes are good for 6 months in freezer (-15°C or below). Thawed working concentrations are good for 2 weeks refrigerated (2-8°C).

The exogenous internal control (1:100 concentration) is stored at -15°C or below. Prepare working stock by adding 990 μ L molecular grade H₂O to single IC tube (containing 10 μ L). The working stock can be stored at 2-8°C. On day of use, dilute working stock an additional 1:100. The final concentration of the final product is 1:100,000. See Appendix B for Internal Control Plasmid info.

4. Supplies / Materials

- Isopropanol, 70%
- RNase AWAY®
- Sterile scrub brushes
- Sterile oyster knives
- Sterile wide mouth containers (1 liter capacity)
- Oyster shucking block
- Chain-mail glove
- Dishwashing gloves
- Serological pipettes (1ml, 10ml, 25ml)
- Filtered pipette tips
- Sterile dilution bottles
- Microcentrifuge tubes (1.7mL)
- 384 well PCR plates; Thermo Fisher Cat. # 4326270
- Optical covers
- MagNAPure 96 Cartridge seals; Roche Cat. # 06241638001
- MagNAPure 96 Processing Cartridge; Roche Cat. # 06241603001
- MagNaPure 96 Output plate; Roche Cat. # 06241611001
- MagNAPure 96 System Fluid; Roche Cat. # 06640729001
- MagNAPure 96 Tips 1000 μ L; Roche Cat. #06241620001

4.1 Bacterial Cultures

- *Vibrio parahaemolyticus* (WA4647 and BAA-240, or equivalent)
- *Vibrio vulnificus* (ATCC 29307 or equivalent)

5. Equipment

- Non-mercury glass thermometer 0°C-10°C
- High Speed blender
- Balance (± 0.1 g)
- Sterile blender jars
- Timer
- Vortex mixer
- Incubator ($35^{\circ}\text{C} \pm 0.5$)
- Refrigerator, $2-8^{\circ}\text{C}$
- Freezer, -15°C to -25°C
- Biological safety cabinets (BSC) or Air Clean PCR stations
- Pipettes P-1000, P-200, P-20
- Multi-channel Pipette (8) $2\mu\text{L}-25\mu\text{L}$
- Applied Biosystems® QuantStudio™ Dx™ Real-Time PCR station
- Roche MagNAPure 96 DNA purification system
- PCR plate centrifuge.

6. Safety Precautions

Vibrio species are pathogenic and should be handled following PHL safety guidelines and assay risk assessment.

UV light can seriously burn skin and eyes. Keep safety shield lowered when UV light is on. Always keep skin covered by lab coat and gloves

7. Specimen Information

Samples are to be shipped properly (adequate ice/cold packs) and temperature maintained between 0 and 10°C upon arrival. Once received and logged-in, the samples are to be placed under refrigeration unless processed immediately.

Samples received over 10°C are considered acceptable only under the following conditions:

The sample is shipped properly (adequate ice/cold packs) and was at an elevated temperature at collection and has had a short transit time (collected and received on the same day).

Sample processing at the lab must be initiated no more than 36 hours after collection. Samples received more than 36 hours after collection are considered unsatisfactory.

8. Quality Control

- Instructions
 - Three process controls are included in every extraction run. To prepare the process control material, enrich *V. parahaemolyticus* WA4647 (*tlh+*, *tdh+*, *trh+*) and *V. vulnificus* ATCC 29307 (*vvha+*) overnight in APW. The overnight enrichment is diluted and a $<10^3$ per ml culture is used as a positive control. The two organisms will also serve as negative

controls for each other. Include an uninoculated APW blank to serve as the uninoculated control. Process control cultures are to be run concurrent with the samples, and accompany the samples throughout incubation, isolation, and confirmation.

- To ensure that all MagNA Pure 96 plastics (i.e. processing cartridge, output plate, internal control tubes, 1000µl tips) and external system fluid are free of interfering contaminants, process controls will serve as quality control.
 - Two amplification controls are included in every PCR run. To prepare the positive PCR control, enrich *Vibrio parahaemolyticus* (ATCC BAA-240, WA4647) and *Vibrio vulnificus* (ATCC 29307) overnight in APW. DNA is extracted individually, combined in equal parts, and divided into 10µL aliquots. Pre-extracted amplification controls should be stored frozen at -15°C or below. Expiration is one year from the date it was prepared. Sterile molecular grade water will be used for the no-template-control (NTC), and will serve as the negative amplification control.
 - Certificates of analysis will be maintained in a binder within the Food lab for all pre-sterilized consumables.
 - Disposable pipettes will be checked for accuracy and tested for sterility.
 - Disposable pipettes (i.e. serological) used to inoculate samples and prepare reagents are checked for accuracy and tested for sterility.
- Frequency
 - Process controls, exogenous internal control, and amplification controls are included with every PCR run.
 - Quality controls will be run on all media and reagents, mastermix, and primer/probe mixes prior to use or concurrent with testing.
 - Certificates of analysis (COA) for each new lot of pre-sterilized consumables will be maintained.
 - Each new lot of disposable pipettes (i.e. serological) will be tested for accuracy and checked for sterility.
- Acceptable Limits
 - See respective media, reagents, mastermix, or primer/probe QC guides for expected or acceptable results.
 - A successful PCR run should meet the following conditions:
 - I. The positive controls should show clear amplification. If NO amplification is present in the positive controls for one or both multiplex's, determine the problem and re-run the sample.
 - II. The negative process control should only show amplification for the exogenous internal control (IC). The no-template-control (NTC) should not show amplification for any of the targets.
 - III. Creeping curves without a logarithmic increase are not considered true amplification. If amplification is present in a negative control, determine the source of contamination, thoroughly clean BSC and pipettes, and discard contaminated reagents and disposables.
 - Results will not be reported without acceptable QC results.
 - All certificates for pre-sterilized consumables will be checked for conformance and initialed by the laboratorian.

- For all disposable pipette (i.e. serological) accuracy checks, a satisfactory accuracy is $\pm 2\%$ of volume tested.
- **Corrective Action**
 - PCR runs for which the NTC is positive or the positive control and/or internal control is negative should be repeated.
 - The Lead Microbiologist should be notified if any run fails QC.
 - If quality controls for mastermix, primers/probes, or disposable pipettes do not meet acceptable criteria, the QC fails and item shall not be used for testing.
 - Media/reagent quality controls not exhibiting the expected growth or reactions will be retested with fresh growth (18-24 hrs) organisms.
 - Samples tested with any failed media/reagent will be considered invalid and will be retested with new media/reagent which has passed quality control.
- **Recording QC Data**
 - Initial quality control data for media/reagents, mastermix, primers/probes, and disposable pipettes will be recorded on its respective QC worksheet. Records are maintained in a binder within the laboratory.
 - Record results for each PCR and Process controls by notating presence (+) or absence (–) on the sample worksheet.

9. Calibration

- **QuantStudio Dx PCR Workstation**

Calibration kits are used to maintain the Real-Time PCR system with 384-Well Block. They include calibration plates to perform a spectral calibration with FAM™, VIC®, ROX™, SYBR® Green, TAMRA™, NED™ dyes, plates to perform region-of-interest (ROI) calibration, normalization calibration, and to run RNase P verification.

Calibration and verification should be run at least every six months and following a Performance Maintenance. Kits are stored at -15°C to -25°C. The maximum degree of accuracy for each dye of interest in fluorescence emission wavelength is $\pm 5\text{nm}$. Follow manufacturer's instruction on performing calibrations.

- **Micropipettor and Thermometers**

Micropipettors are calibrated at appropriate volumes annually and checked for accuracy quarterly. At a minimum quarterly checks are performed at 100%, 50%, and 10% of nominal volume.

Non-mercury glass thermometers will be sent out for annual calibration and checked for accuracy quarterly by a certifying vendor on-site. Long stem digital thermometers will be replaced yearly.

“As Found” and/or “As left” calibration data for micropipettes or thermometers must indicate that the initial calibration or recalibration passed. Acceptable tolerance limits will be obtained as pre-determined by the manufacture or ISO 17025 accredited service by the certifying vendor.

If the “As found” calibration data for micropipettes or thermometers indicate that the calibration or recalibration failed, a PHL Quality Improvement (QI) Form must be filled out for all affected samples.

Calibration/re-calibration certificates for all micropipettes and thermometers will be checked for conformance and initialed by the Supervisor or Lead prior to use. Calibration certificates are maintained in a binder within the laboratory.

10. Procedure

10.1 Sample Accessioning

- a. Samples are collected, transported, and processed in accordance with Recommended Procedures for the Examination of Sea Water and Shellfish described by the American Public Health Associationⁱⁱⁱ.
- b. Oyster samples are removed from the shipping container and the sample submission form is located. At a minimum the sample submission forms must contain the following information: collector’s name, harvest area, sampling station, time and date of collection.
- c. A laboratory testing worksheet is generated for each sample.
- d. Both the sample submission form and the testing worksheet are stamped with the appropriate laboratory number.
- e. One oyster from each bag is opened to take tissue temperature. The temperature is recorded on the sample submission form.
- f. The bag of oysters is labeled with its associated sample ID and placed into a 2-8°C refrigerator unless processed immediately.

10.2 Sample Preparation- Scrubbing

- a. The intent of the assay is to determine the concentration of V_p and V_v in the oyster tissue and liquor. Any material on the outside of the oyster that gets introduced into the interior of the animal during shucking can alter the concentration.
- b. The sink must be clean before scrubbing can begin. Wash the sink with water or soap and water.
- c. The gloved hands of the analyst are to be washed with soap immediately prior to cleaning the shells of debris. The gloves worn are latex, nitrile and/or stainless steel mesh to protect analyst’s hands from injury.
- d. Using sterile scrub brushes, each oyster is cleaned under cold running water. All barnacles, mud, vegetation and debris should be removed.
Note: The faucet used for rinsing the shellfish should not contain an aerator. Pay close attention to the hinge and shell seam. A sterile brush should only be used for one sample. Do not re-use brushes when scrubbing multiple samples. Any oyster that does not tightly close during handling is likely dead and should be discarded. In addition, any oyster whose shell is broken to expose tissue should be discarded.
- e. A representative sample of at least 12 shellfish is used for analysis.
- f. After cleaning each oyster place the animal upside down on a clean paper towel lined tray. Ensure that you have labeled the tray with corresponding sample number.

Laying the oysters upside down will prevent the liquor (fluid inside a closed oyster) from draining out of the oysters while waiting to be shucked. Clean trays must be used for each sample.

- g. Once cleaned, return the oysters to the refrigerator to dry or towel dry them for immediate shucking.

10.3 Sample Preparation- Shucking

- a. In order to accurately quantify V_p and V_v in oyster tissue it is very important to avoid introduction of bacteria (V_p or other) into the oyster tissue.
- b. The sink must be disinfected before shucking can begin. Wash the sink with water or soap and water. Completely dry the sink. Wipe the sink down with 70% isopropanol and allow it to air dry.
- c. Place a sterile pre-weighed tissue collection container on the sink counter.
- d. Disinfect a shucking block by washing with soap and water, and wiping down with 70% isopropanol. Place block on the sink counter to air dry.
- e. Place the oyster sample to be shucked on the sink counter.
- f. Put on clean nitrile gloves.
- g. Over one of the gloves put on a chain-mail glove. The chain-mail glove should be on the hand that will not be holding the knife.
- h. Put another nitrile glove on over the chain-mail glove. Cover both hands in 70% isopropanol and allow them to air dry.
- i. Grab and hold each oyster with the chain-mail hand and use the other hand and a sterile oyster knife to shuck each oyster.
- j. A fresh knife, shucking container and gloves must be used for each sample.
- k. Use the disinfected shucking block while shucking to minimize knife accidents and to protect the counter surface.
- l. Collect all tissue and liquor (fluid) in the sterile pre-weighed container. 10^{-1}
- m. The shucking block and counter must be washed and sterilized between samples.

10.4 Sample Processing- Setting up MPN

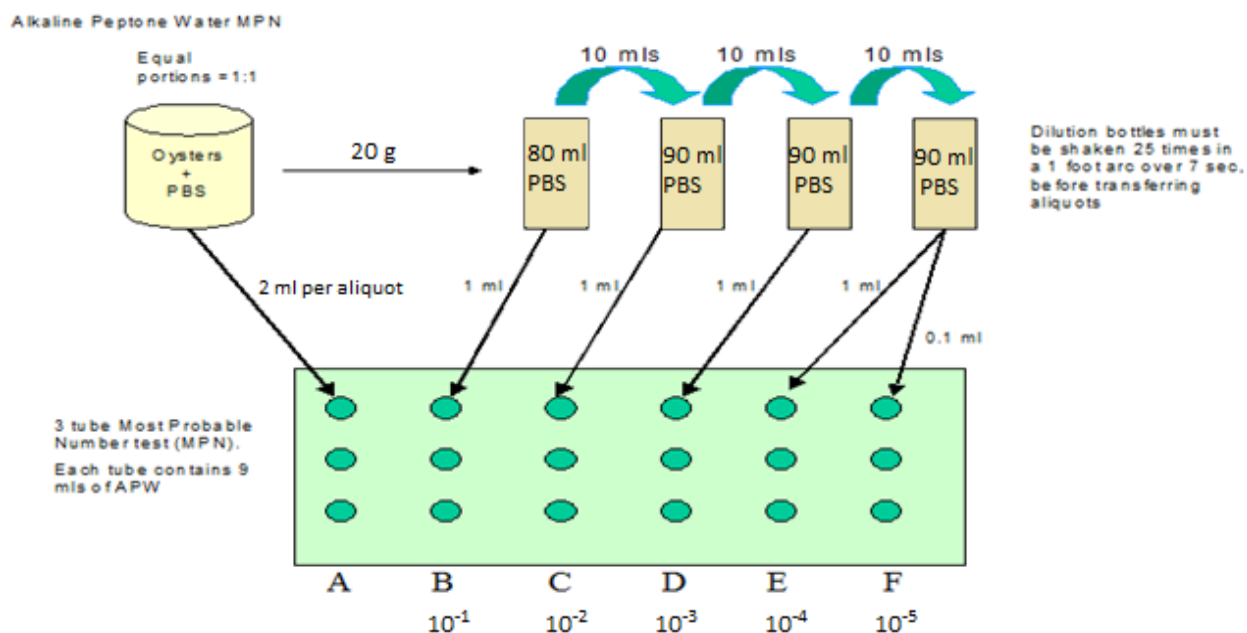
Enumeration in this assay is achieved by conducting an MPN (most probable number) analysis.

- a. Weigh the shellfish tissue collection container. Subtract the original container weight from the total weight to determine the amount of oyster tissue and liquor.
- b. Record the weight of tissue on the sample worksheet.
- c. Transfer the oyster tissue and liquor to a sterile blender jar.
- d. Add an equal weight of diluent (PBS) to the sample container. If needed, PBS can be used to rinse any residual tissue from the container just as long as a 1:1 dilution can be maintained ($\pm 0.1g$).
- e. Transfer the PBS to the blender jar. Record the weight of PBS used on the sample worksheet.
- f. Blend the shellfish sample with PBS at high speed for 90 seconds (60 to 120 seconds is acceptable).

The resulting homogenate should be relatively smooth. If the blender isn't generating a smooth homogenate, it is advisable to service the blender (replace blades).

- g. From this homogenized sample, set up a 3-tube most probable number (MPN) serial dilution series. Use PBS for making dilutions and alkaline peptone water (APW) as the enrichment broth in each of the MPN tubes. See Figure below.

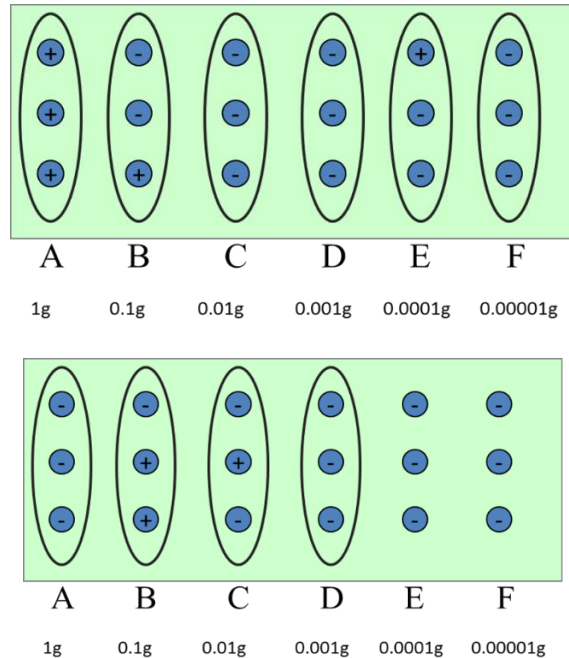
The initial 1:10 dilution is prepared gravimetrically with PBS (20 g of sample homogenate to 80 mL of PBS). Record the weight and volume used on the sample worksheet for the sample worksheet. All successive dilutions are prepared volumetrically.



- h. Incubate at $35^{\circ}\text{C} \pm 1$ for 18 to 24 hours. Write sample number, date, time, and analyst initial on the A1 tube, prior to placing in incubator.

10.5 Reading MPN

- a. Each APW tube must be checked for growth following 18-24 hrs. of incubation. Use the following criteria to select tubes for further testing.
- Examine all tubes for turbidity. Examine each tube with a light source shining through the tube.
 - Record all positive and negative results on the lab worksheet.
 - The following examples illustrate the selection process. Each tube is labeled as +/- for turbidity. The dilutions circled should be selected for further testing.



- If one tube in a given three tube dilution series is positive for turbidity, all tubes in that series must be tested (you will still record the actual positive or negative values).
- In addition, test one complete dilution series beyond the last series that contained any growth and all tubes of higher concentration.

10.6 DNA Extraction

DNA extraction must be initiated within the 18-24 hour incubation window.

Reagent prep should be carried out in the Pre-PCR room in order to minimize the potential for contamination. Once extraction is complete, the material is considered stable and may be stored at 2-8°C for 1-2 days or frozen at -20°C to -80°C for extended periods. NOTE: Multiple freeze thaw cycles should be avoided due to potential DNA degradation.

- Due to the high number of sample tubes it is necessary to create a document to track the location of each one. This document is referred to as the "MagNA Pure Plate Map".
- After the MagNA Pure Plate Map is created, load the MagNA Pure 96 cartridge accordingly. 200µL of each sample should be added to the 96 well cartridge. Include 200µL of Vp culture, 200µL of Vv culture, and 200µL of uninoculated APW. These will act as process controls for the assay.
- Once loaded, seal the MagNA Pure cartridge with an adhesive cartridge seal.
- Label the cartridge with the sample number, date, initials and label as "Pre-Extract". The specimen can now be loaded onto the MagNA Pure 96 instrument.
- Confirm that the MagNA Pure 96 instrument and its linked computer are turned on.
- Ensure that the correct MagNA Pure kit is selected "DNA/Viral SV 2.0".
- Select the protocol "Pathogen Universal 200.3.1"
- Sample volume should be entered as 200µL.
- Elution volume should be entered as 100µL.

- j. Next to the Internal Control section, click the More Options icon. Scan the barcode located on the IC tube. Enter the number of tests. This will determine the amount of IC needed. Since IC is prepped for single use, the auto-filled expiration date is not relevant.
- k. Enter in your sample order. Be sure that the correct cartridge wells are highlighted on the screen. Incorrect set up here will lead to a failed extraction.
- l. Click the “Stage Set-up” button.
- m. Begin adding in the appropriate reagents/plastics in accordance with the outlined requirements on the MagNA Pure load screen (software).
- n. Once the reagent trays are completely loaded and the tips are adequately filled, place the remaining trays back into the instrument.
- o. Remove the cartridge seal from the processing cartridge and place into the instrument. Discard the seal into an autoclave waste container.

- p. Ensure that all plastics, reagents and sample cartridges are in place and accounted for on the computer screen.
- q. Close the door and press the “start extraction” button.
- r. Note the time that the run will be completed. The final extracted template DNA will be refrigerated on-board the MagNA Pure 96 instrument until it is removed. It is however not advisable to leave the extract uncovered for any length of time.
- s. Once completed, open the door, remove the extracted DNA, immediately seal the cartridge with a new cartridge seal, and refrigerate at 2-8°C until ready for PCR (if PCR is to be completed in 1-2 days). If PCR will not be complete in the next two days freeze the DNA at -20°C to -80°C.

10.7 **MagNAPure 96 Waste Removal and Decontamination.**

- a. Remove all soiled plastics, replace used tips, and wipe the trays with 10% bleach, isopropyl alcohol and RNase Away using Manufacturer’s suggested cleaning procedure. Run the UV decontamination protocol.
- b. If waste bottle is full, follow MagNAPure 96 Waste Disposal Procedure listed below.
 - I. Instrument will indicate waste container is full.
 - II. Attach empty waste container to instrument and secure caps on full container.
 - III. Carry full waste container with caps closed to the sink, set inside sink, and remove small cap.
 - IV. Tip container onto side with small cap and allow to drain into sink. As it drains, you may need to tilt the container to ensure complete drainage of liquids.
 - V. Spray sink and container with 70% Isopropanol.
 - VI. Pour entire bottle of 70% Isopropanol (~ 500ml) into container, secure cap, and carefully invert to mix.
 - VII. Allow Isopropanol to sit for 10 min.
 - VIII. Spray sink and container again with 70% Isopropanol and wipe down container. Pour Isopropanol from inside container into the sink and secure caps.
 - IX. Rinse sink and exterior of container with water.
 - X. Spray exterior of container with 10% bleach, allow 3 minute contact time, and then rinse with water.
 - XI. Container can be stored in lab with secured caps until next use.
- c. For routine MagNA Pure 96 maintenance, follow the MagNA Pure 96 Daily Maintenance Log (for start-of-day and end-of-day instructions) and MagNA Pure 96 Post Run Cleaning Log

instructions. Forms can be found in the Master Document Control or link to the following address:

P:\EHSPHL\PHL\MICRO\COMMON\ENTERICS - FOOD\QC\Media QC\501.4206.docx

P:\EHSPHL\PHL\MICRO\COMMON\ENTERICS - FOOD\QC\Media QC\501.4207.docx

10.8 PCR Mastermix Preparation

Mastermix preparation is performed in the Pre-PCR room, within an Airclean hood. This includes primer and probe manipulations and mastermix loading onto the PCR plate. Thorough decontamination before and after use of the Airclean hood is advisable.

Note: A person who has previously in the same day worked with amplicon should not re-enter the Pre-PCR lab.

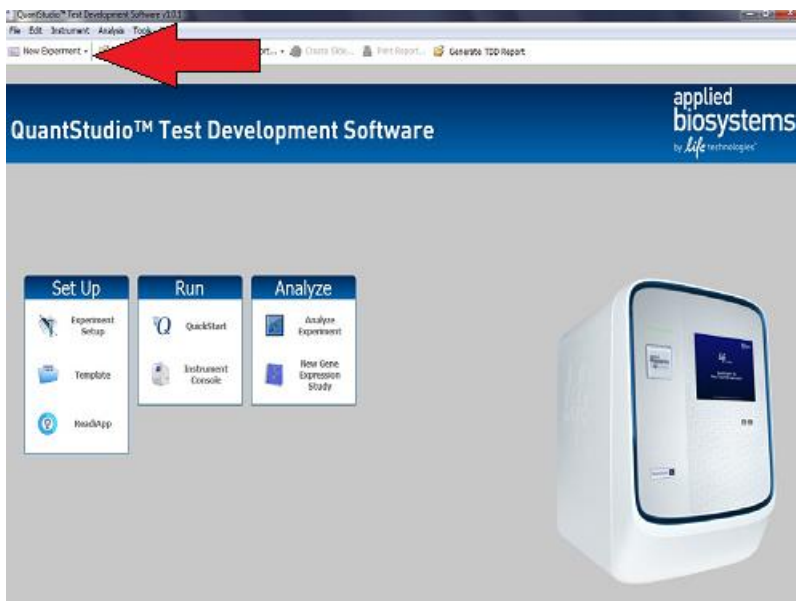
- a. Prepare a PCR platemap using the MasterMix Prep worksheet.
(P:\EHSPHL\PHL\MICRO\FOODLAB\Vibrio\Master Mix Documents\Mastermix Template).
Be sure to include positive and negative process controls, in addition to a positive and negative amplification control.
- b. Using the worksheet determine the number of reactions needed. This will help you determine how much of each mastermix component will be required. It is advisable to prepare several reactions more than are needed to account for pipetting variability.
- c. Once in the Pre-PCR room, follow mastermix recipe and MagNAPure plate map to prepare Multiplex 1 and 2. The mastermix can be prepared in a microcentrifuge tube or sterile disposable reagent reservoir.
- d. Briefly vortex (swirl, pipette up and down, or equivalent) to completely mix the components.
- e. Using a pipette (multichannel advisable), add 18 μ L of mastermix to each appropriate well (384 well plate) according to the PCR platemap.
- f. Once the 384-well plate is loaded with mastermix, cover the plate with aluminum foil, place the plate in a biological transport container (sealed box), and transport to an available AirClean hood within the Food laboratory (Alternatively the Template Addition Room can be used).

10.9 Template Addition

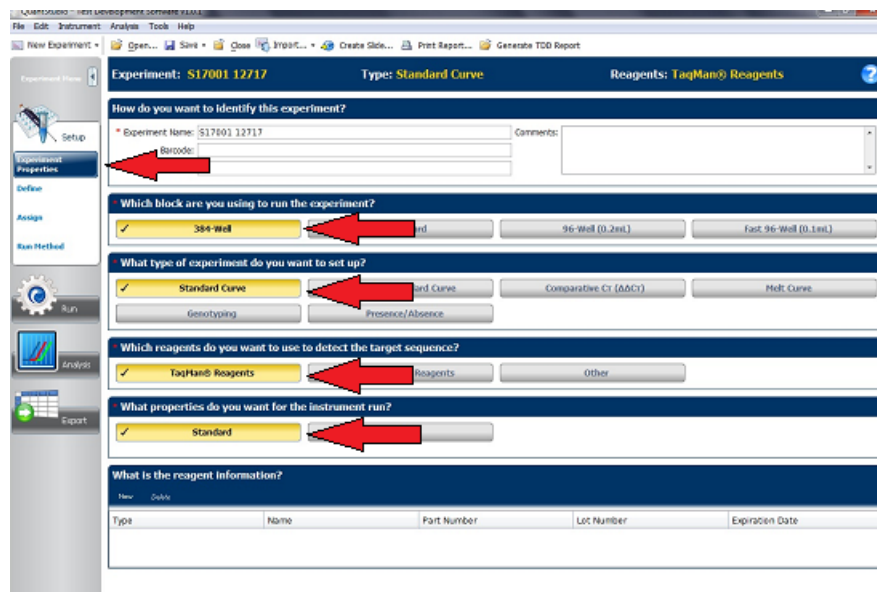
- a. Add the extracted DNA template to the appropriate wells according to your PCR platemap. Use 2 μ L of DNA for a total reaction volume of 20 μ L.
- b. Once all wells are loaded including the positive and negative amplification controls, seal the plate with an optical seal.
Avoid touching either side of the seal as the adhesive side will come into contact with your reactions (poses a contamination risk) and the outside must be clear of smudges to allow accurate readings. Apply the seal using the plastic applicator supplied with the instrument. Take care to completely seal each well. Any unsealed well will rapidly evaporate during PCR and lead to inaccurate results.
- c. Centrifuge the plate briefly to remove bubbles from the wells and ensure that the template is in contact with the reaction mix.

10.10 Setting up Real-Time PCR Station

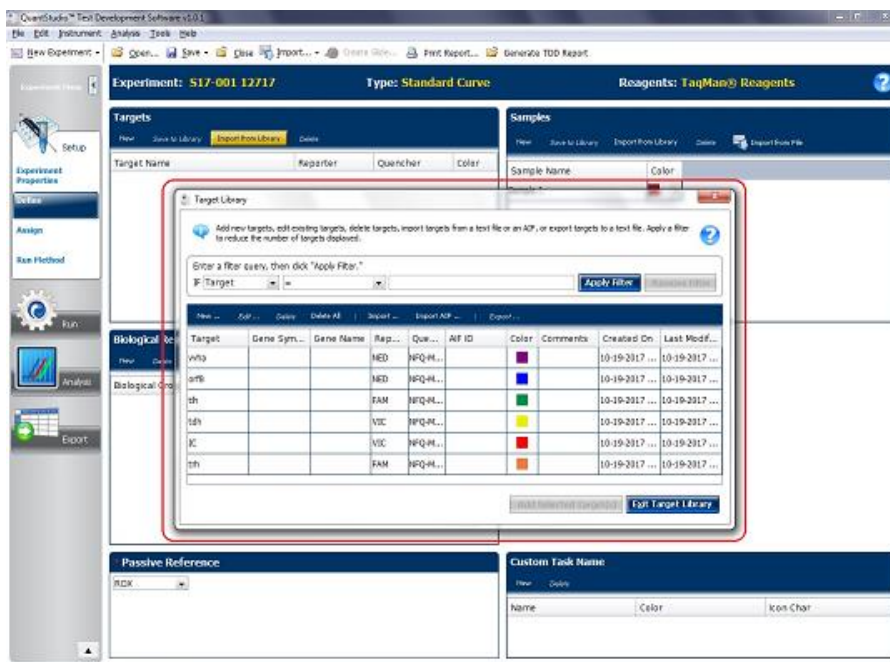
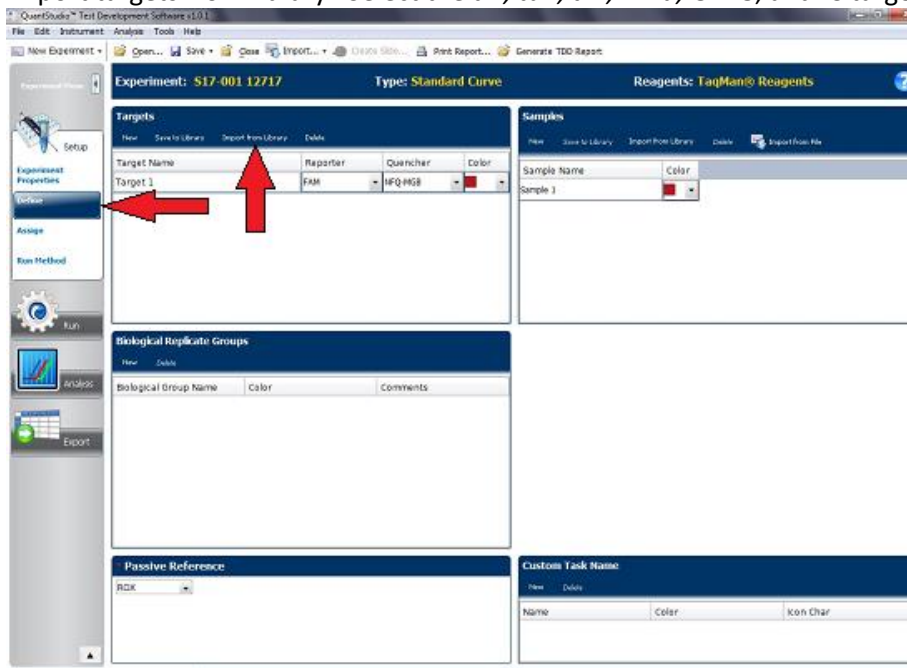
- a. Turn on the Applied Biosystems® QuantStudio™ Dx™ instrument and the computer.
- b. Open the Test Development software, under the File menu select “New Experiment”.



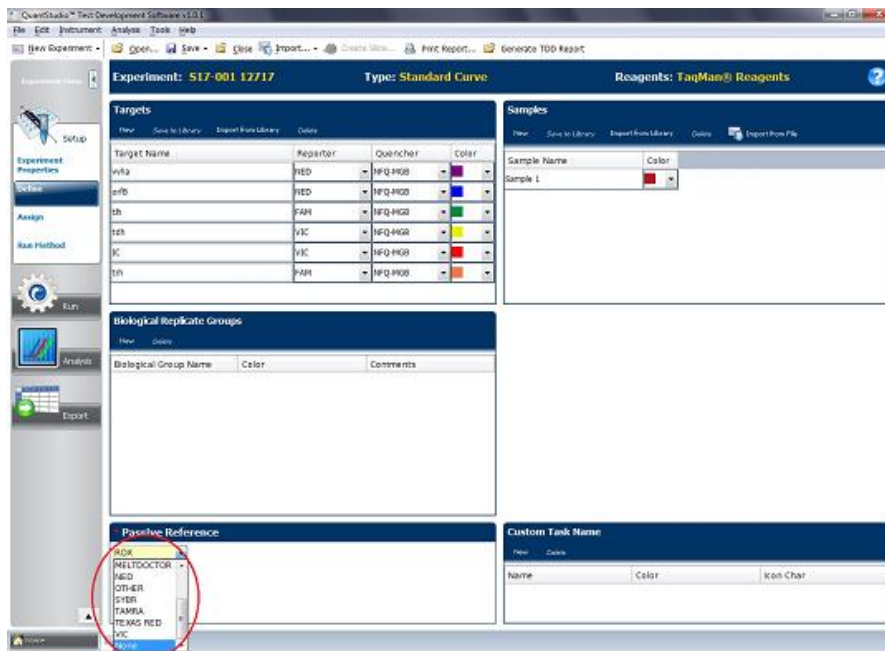
- c. Rename experiment with the appropriate sample numbers and date of run.
- d. Under the tab “Experiment Properties” ensure “384-Well Block”, “Standard Curve”, TaqMan® Reagents”, and “Standard” (for run mode) are selected.



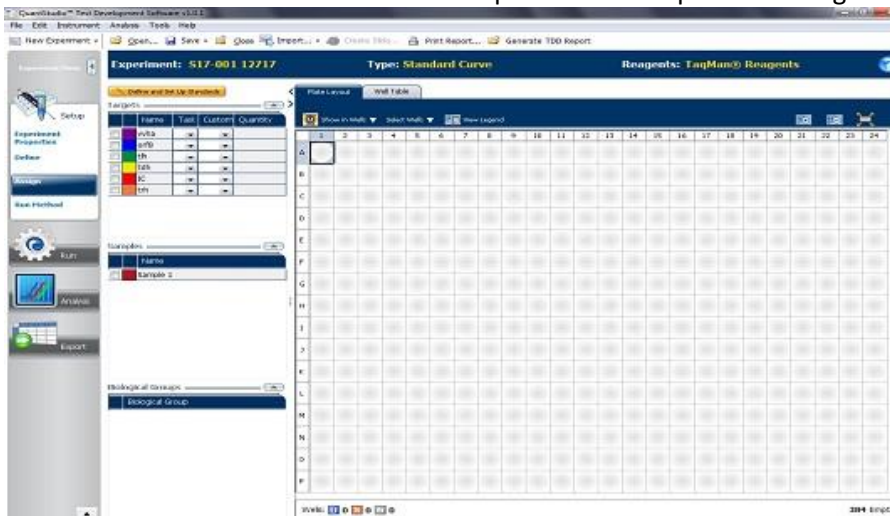
- e. The “Define” tab is used to select the targets of detection. The targets are saved to the library. Import targets from library. Select the *tlh*, *tdh*, *trh*, *vvha*, ORF8, and IC targets.



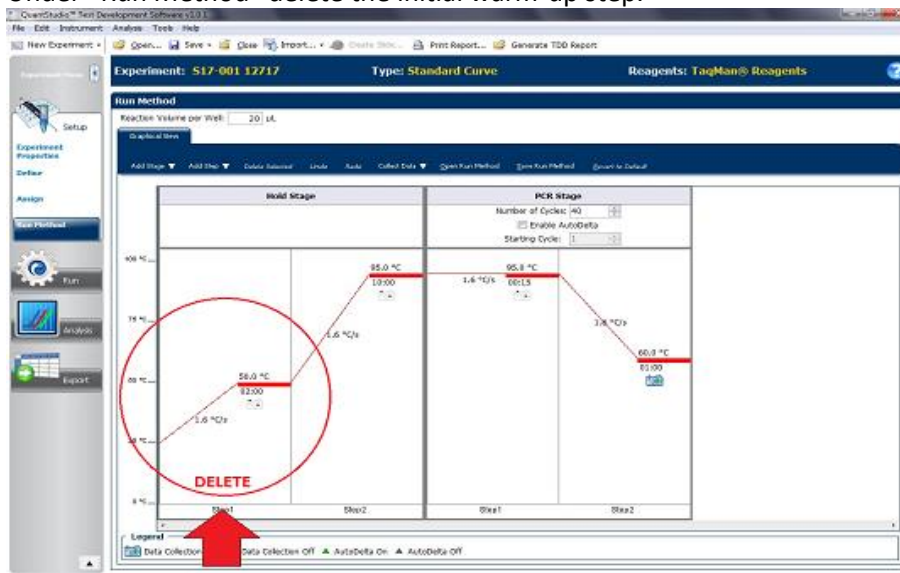
- f. At the bottom of the screen, select “ROX” from the drop down menu regarding “Passive Reference”



g. Select the "Assign" tab and assign the appropriate wells with the corresponding targets of interest. Be sure to double check the map and 384-well plate are in agreement.

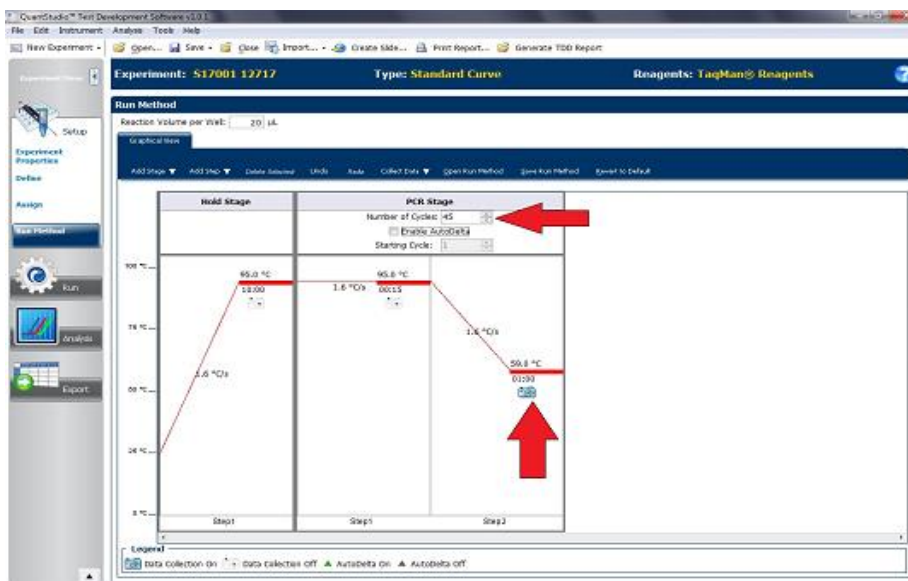


h. Under “Run Method” delete the initial warm-up step.



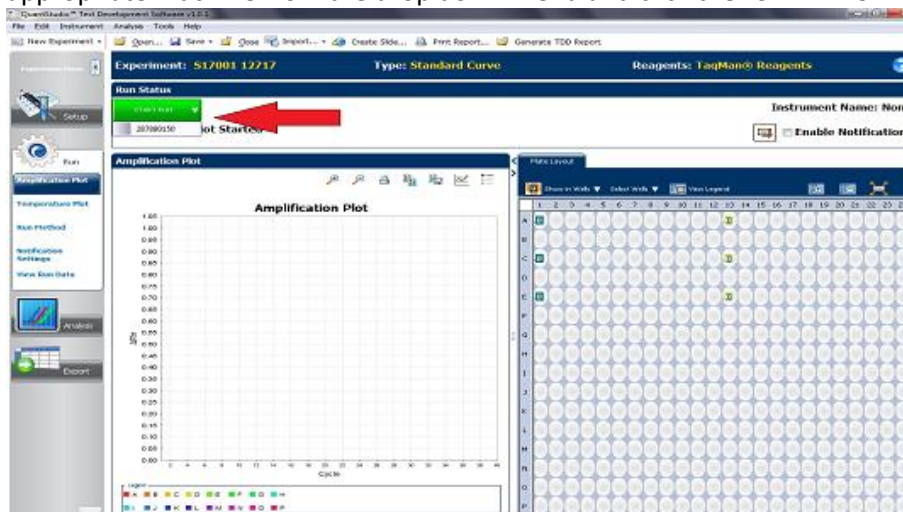
i. After doing so the parameters are the following:

- 95° C for 10 minutes
- 40 cycles
- 95° C for 15 seconds
- 59° C for 60 seconds



- j. Turn on the instrument user interface by touching the touchscreen. To open loading tray, touch the eject button.
- k. Load the plate and close tray.

- I. From the PC, click the “Run” tab and find the green “START RUN” button. Select the appropriate machine from the drop down menu and click the “START RUN” button.



- m. Save run file using sample numbers and date.

11. Waste Management

The biological material is rendered non-hazardous through use of the MagNA Pure 96 System and associated reagentsⁱⁱ. The reference refers to the MagNA Pure Compact System, however; all of these instruments share the same buffer system, process, and concentrations. The remaining waste is considered flammable by Department of Ecology Standards and will be collected as hazardous waste for disposal.

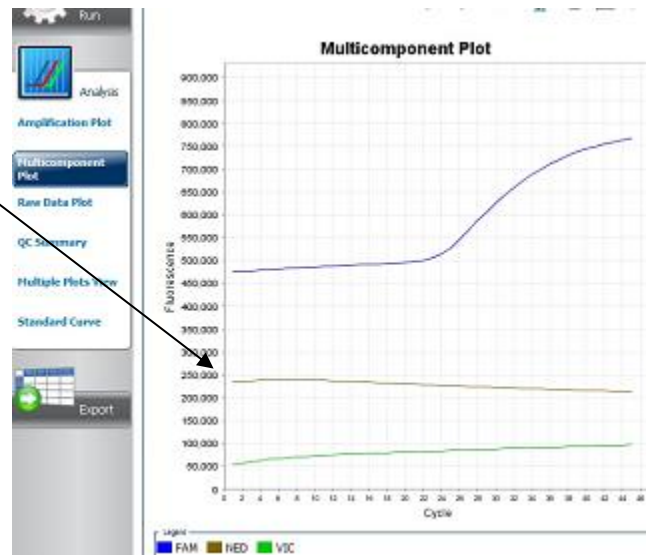
All other plastics and glassware containing the sample will be treated by autoclaving using appropriate conditions.

Date of disposal of each sample is recorded on the respective sample worksheet.

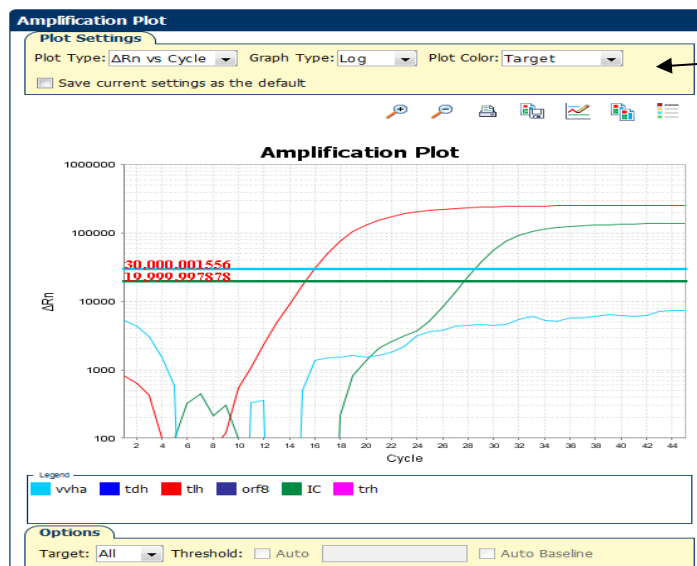
12. Interpretation

- a. Once run is complete, remove plate from instrument and discard in appropriate waste receptacle.
- b. Select “Analysis Settings” on the Amplification Plot screen.
- c. Change all thresholds and baseline settings to “manual” and set all thresholds, except trh, to 0.04 and leave baseline settings at 5 to 15. Set trh threshold to 0.08.
- d. Apply analysis settings and exit to Amplification Plot screen.
- e. Record quality control Ct values on the *Vibrio parahaemolyticus* Real-Time PCR Ct Value QC worksheet. For all targets record Ct value from the positive mastermix control. Record Ct value for the Internal Control (IC) using the negative mastermix control. All Ct values should be recorded with the threshold set at 0.04 and baseline set at 5 to 15.
- f. Select all wells in the plate by clicking in the upper left box of the plate layout.
- g. View each target individually and make necessary changes to the threshold and baseline. The threshold should be set above background levels. It may be necessary to change baseline settings to lower background levels.
- h. If baseline changes are necessary - view individual wells in the Multicomponent Plot screen. Change baseline settings as needed to exclude early background noise.

Exclude early background noise when setting baseline



- i. Once threshold and baseline are set at appropriate levels, record results from each well for every target. Targets within the amplification plot may have high background in the early stages of the run (i.e. <10 cycles). Disregard background that crosses the threshold before cycle 10. Change the Plot color to "Target" to help read results.



Plot Color = Target

Positive = amplification above threshold
Negative = No amplification

12.1 Procedure for Abnormal Results

If abnormal results appear to be caused by cross contamination (i.e. late CT value) rerun real-time PCR in duplicate of suspected contaminated wells

If duplicate results are in agreement, report these results. If the duplicates differ, report the result that is in agreement with the original qPCR run.

Positive pathogenic markers (*tdh*, *trh*, ORF8) in absence of *Vibrio parahaemolyticus* marker (*tlh*)

- I. Current findings do not support pathogenic markers being present without *Vp* being present. Real-time PCR reactions resulting in this situation should be re-run upon Lead Microbiologist discretion.
- II. The presence of the *trh* gene in the absence of the *tlh* gene has been documentedⁱ. This is due to the *Vibrio parahaemolyticus trh* gene having 98% homology with the *trh* gene of *Vibrio alginolyticus*ⁱ. Any *trh* positive wells must be *tlh* positive as well.

12.2 Interfering Substances

Vibrio alginolyticus possesses a *trh* gene with 98% homology to the *trh* gene in *Vibrio parahaemolyticus*ⁱ. Most probable number values for *trh* should be reported only if *tlh* is present in the corresponding tube. Tubes only positive for *trh* should not be accounted for when generating the MPN value.

13. Calculations

Upon determination of positive reactions, record the number of confirmed positive tubes per dilution series onto the *Vibrio* Sample Worksheet, and generate an MPN index. MPN values (concentration) of each target is derived from the FDA Bacteriological Analytical Manual (BAM) MPN Calculator. This Excel document can be located in Appendix 2 of the FDA BAM. To compute an MPN value, follow the instructions as noted in the FDA BAM MPN Calculator.

Unusual MPN indexes are typically due to contamination. It may be necessary to re-extract and/or re-run PCR. If this does not resolve the issue, further investigation is required to determine the source of contamination.

14. Reference Range

Reportable Range

tlh: <0.36 MPN/g to >110,000 MPN/g

tdh: <0.36 MPN/g to >110,000 MPN/g

trh: <0.36 MPN/g to >110,000 MPN/g

ORF8: <0.36 MPN/g to >110,000 MPN/g

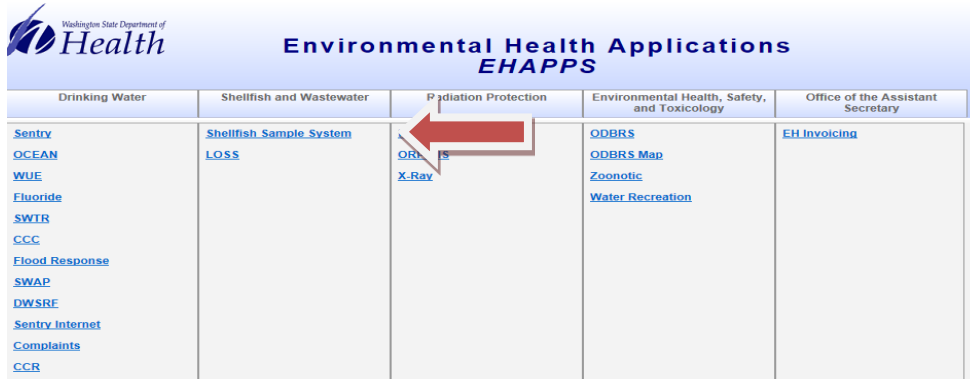
vvhA: <0.36 MPN/g to >110,000 MPN/g

15. Reporting Results

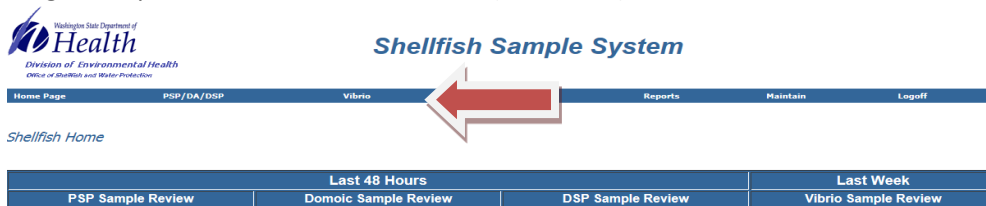
15.1 Environmental Health Applications (EHAPPS) database

Access to the database must be authorized. Lead Microbiologist or Supervisor will facilitate the authorization process.

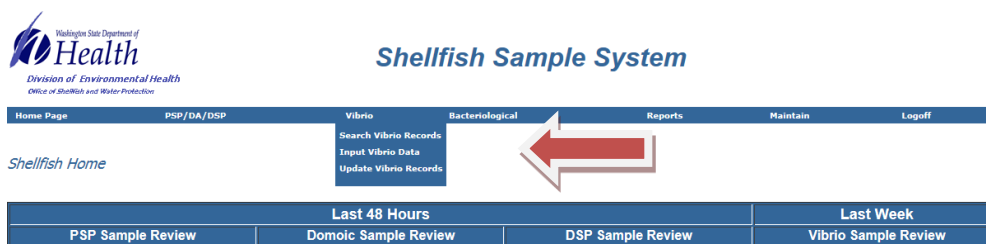
- a. After entering web address, find column “Shellfish and Wastewater” and click “Shellfish Sample System” (red arrow)



- b. Along the top, hover mouse over “Vibrio” (red arrow)



- c. Click “Input Vibrio Data” (red arrow)



- d. The sample number will auto-populate. Enter all information and data provided from Vibrio Sample Submission form. Enter final test results when available.

The screenshot shows the 'Input Vibrio Data' form. It is divided into two main sections: 'Shellfish Data' and 'Lab Data'.
Shellfish Data: Includes fields for 'PreviousYear' (radio button), 'CurrentYear' (radio button), 'Vibrioid' (text input: 2014 00001), 'Date Collected' (calendar), 'Low Tide Time' (checkbox), 'Time Collected' (AM/PM), 'Organization' (dropdown), 'Water Temp (-3ft)' (text input), 'Cert#', 'Surface Water Temp', 'Site Name', 'Shore Water Temp', 'Site ID', 'Tissue Temp', 'Sample Type', 'Shell / Shucked', 'Species', 'Fresh / Frozen', 'No. of Organisms', 'Ambient Air Temp', 'Sampler', and 'Select Harvest Conditions' (checkboxes for Overcast, Rainy, Sunny, Windy). There is also a 'Sample Comments' text area.
Lab Data: Includes fields for 'Sample Weight' (text input), 'Salinity' (text input), 'Vibrio parahaemolyticus' (text input), 'TLH MPN' (dropdown), 'TDH MPN' (dropdown), 'Vibrio Vulnificus' (dropdown), 'CFU/g' (text input), 'Date Received' (calendar), 'Time Received' (AM/PM), 'Date Examined' (calendar), 'Time Examined' (AM/PM), 'Date Reported' (calendar), 'Time Reported' (AM/PM), 'Shellfish Tissue Temperature at Lab' (text input), 'Exception' (dropdown), and 'Lab Comments' (text area).
 At the bottom, there are 'Required' (checkbox), 'Save', 'Reset', and 'Cancel' buttons.

15.2 Notification of Test Results

a. Environmental Health Applications (EHAPPS)

All test results will be entered into the Shellfish Sample System via EHAPPS. Results are reviewed and checked off by the Lead Microbiologist.

b. Email

Test results can be emailed to the Office of Shellfish and Water Protection (OSWP) after Lead Microbiologist approval and signature.

c. Phone

For STAT results (per request of OSWP), the Lead Microbiologist will contact the appropriate personnel at OSWP.

15.3 Archiving Results & Retention

a. Filing Results

The Vibrio Testing Worksheet and Sample Submission Form are to be filled in a filing cabinet located within the Food and Shellfish Bacteriology Laboratory.

All other documents (i.e. Mastermix worksheet, PCR Plate Map, MP96 Plate Map, Sample Tracking worksheet, etc.) are to be scanned and uploaded into the Scanned Testing Documents folder under the appropriate year on the PHL P: Drive server. All scanned documents for a given day can be saved under this folder as the date (MMDDYY).

For example:

P:\EHSPHL\PHL\MICRO\FOODLAB\Vibrio\2019 Vp Season\Scanned Testing Documents\060119

All uploaded testing documents will be reviewed by the Lead Microbiologist prior to discarding any hard copies.

b. Retention

Reports and results for samples tested will be archived according to the Department of Health Records Retention Schedule.

EHAPPS database is maintained by the Office of Shellfish and Water Protection.

16. References

ⁱ González-Escalona, Narjol, George M. Blackstone, and Angelo DePaola. Characterization of a *Vibrio alginolyticus* strain, isolated from Alaskan oysters, carrying a hemolysin gene similar to the thermostable direct hemolysin-related hemolysin gene (*trh*) of *Vibrio parahaemolyticus*." *Applied and environmental microbiology* 72.12 (2006): 7925-7929.

ⁱⁱDauphin L. A. et. al. Evaluation of Automated and Manual Commercial DNA Extraction Methods for Recovery of *Brucella* DNA from suspensions and spiked swabs. 2009. *JCM* Vol. 47, No. 12. p. 3920-3926.

ⁱⁱⁱThe American Public Health Association, Inc. *Recommended Procedures for the Examination of Sea Water and Shellfish*. 4th ed., 1970.

17. Appendix

Appendix A- Primers and Probes

VIBRIO PARAHAEMOLYTICUS

TRH

Primers

Trh 627F

ATA CCT TTT CCT TCT CCW GGT TC

Trh 731b R

TTG TCC AGT AGT CAT CAA CGA TTG

Trh Glov R

TTG TCC AAT AGT CCT CCA CAA TTG

(Ward) Probe-- Trh P

FAM TAT TTG TYG TTA GAA ATA CAA CAA T **MGBNFQ**

(WA PHL *Vibrio* Internal Control)

Primers

WA IC F

GGC GAA GCG AAT CTG GAA A

WA IC R

GGT GTA GTT GTG CGT GTA ATA TGA GA

Probe-- WA PHL ICP

VIC CGT AAG ACA ATC TGA TAG TAG T **MGBNFQ**

Orf8

Primers

Orf8 F TCA CCT GAG GAC GCA GTT ACG

Orf8 R TTC AAT TGT AGA ACC GCC AGC TA

Orf8 Probe

NED_ TCC TGC TGT ACT TTT AG **MGBNFQ**

TLH (69 bp amplicon)**Primers****Tlh-F** CCG CTG ACA ATC GCT TCT C**Tlh-R** TTT GAT CTG GCT GCA TTG CT**Tlh probe****FAM** ACC ACA CGA TCT GGA GCA ACG ACG **MGBNFQ****TDH (94 bp amplicon)****Primers****TDH-F 2013** TAT CCA TGT TGG CTG CAT TC**TDH-R 2013** CGA ACA ACA AAC AAT ATC TCA TCA GA**TDH Probe****VIC** TGT CCC TTT TCC TGC CCC CGG MGBNFQ***VIBRIO VULNIFICUS*****VVHA (79 bp amplicon)****vvha-F** GAT CGT TGT TTG ACC GTA AAC G**vvha-R** TGC TAA GTT CGC ACC ACA CTG T**vvha Probe****NED-CAA AAC GCT CAC AGT CG-MGB probe****Appendix B- Internal Control Plasmid**

The *Vibrio parahaemolyticus* and *Vibrio vulnificus* enumeration and detection through MPN and real-time PCR assay utilizes an exogenous internal control (WA IC). This plasmid is added to the during DNA extraction for the detection of matrix inhibition or other assay failures. The 73bp fragment can be synthesized and clones into a pIDTSMART-AMP plasmid by Intergrated DNA Technologies (IDT), Ref ID: 88772700.

Sequence:

GGCGAAGCGAATCTGGAAAACGTAAGACAATCTGATAGTAGTATATTTCTCATATTACACGCACAACACTAC
ACC

Additionally, the Invitrogen OneShot Top10 Chemically Competent Cells and QIAGEN Plasmid Midi Kit can be used collectively to manufacture and purify additional plasmid DNA.

Protocol for Transforming Chemically Competent Cells.

This section provides a procedure to transform Invitrogen One Shot TOP10 chemically competent E.coli via regular Chemical transformation protocol, as described by the manufacturer's instructions.

Step-by-step Procedure:

1. Prepare a plasmid dilution by transferring 1 μ l of 40mM Plasmid (stock) into 3 μ l of molecular water. Briefly place on ice.
2. Thaw, on ice, one 50 μ L vial of One Shot[®] cells for each ligation/transformation.
3. Pipet 1 μ l of plasmid dilution (from step 1) directly into the vial of competent cells and mix by tapping gently. Do not mix by pipetting up and down. The remaining ligation mixture(s) can be stored at -20°C .
4. Incubate the vial(s) on ice for 30 minutes.
5. Incubate for exactly 30 seconds in the 42°C water bath. Do not mix or shake.
6. Remove vial(s) from the 42°C bath and place them on ice.
7. Add 250 μ L of pre-warmed (room temperature) S.O.C medium to each vial. S.O.C is a rich medium; sterile technique must be practiced to avoid contamination.
8. Place the vial(s) in a microcentrifuge rack on its side and secure with tape to avoid loss of the vial(s). Shake the vial(s) at 37°C for exactly 1 hour at 225 rpm in a shaking incubator.
9. Spread 100 μ L from each transformation vial on separate, labeled LB agar plates. The remaining transformation mix may be stored at 4°C and plated out the next day, if desired.
10. Invert the plate(s) and incubate at 37°C overnight.
11. Select isolated colonies and pick to LB medium with ampicillin.

Protocol for Plasmid DNA Purification using QIAGEN Plasmid Midi Kit

This protocol is designed for preparation of up to 100 μ g of high or low copy plasmid DNA using the QIAGEN Plasmid Midi Kit. Consult the manufactures instructions for additional information.

Before starting:

Prepare Buffer P1 according to step 5

Prepare Buffer P2

Prepare buffer P3

Step-by-step Procedure:

1. Pick up to 2 colonies per plate from a freshly streaked selective plate and inoculate a starter culture of 5 ml LB medium containing the appropriate selective antibiotic. Incubate for approx. 8 hr. at 37°C with vigorous shaking (approx. 300 rpm).

Use a tube or flask with a volume of at least 4 times the volume of the culture.

2. Dilute the starter culture 1/500 to 1/1000 into selective LB medium. For high-copy plasmids, in a 250ml flask inoculate 25 ml medium with 50 µl of starter 8hr culture. Grow at 37°C for 12–16 h with vigorous shaking (approx. 300 rpm). Either change shaking incubator platform or attach flask holder to allow for vigorous shaking.

Use a flask or vessel with a volume of at least 4 times the volume of the culture. The culture should reach a cell density of approximately $3-4 \times 10^9$ cells per milliliter, which typically corresponds to a pellet wet weight of approximately 3 g/liter medium.

3. Harvest the bacterial cells by centrifugation at 6000 x g for 15 min at 4°C. Transfer to 50ml centrifuge tubes (falcon or equivalent). Decant the supernatant and retain the pellet.

If you wish to stop the protocol and continue later, freeze the cell pellets at –20°C.

4. Resuspend the bacterial pellet in 4 ml Buffer P1. Vortex in Falcon tubes until no clumps are visible.

For efficient lysis, it is important to use a vessel that is large enough to allow complete mixing of the lysis buffers. Ensure that RNase A has been added to Buffer P1.

If LyseBlue reagent has been added to Buffer P1, vigorously shake the buffer bottle before use to ensure LyseBlue particles are completely resuspended. The bacteria should be resuspended completely by vortexing or pipetting up and down until no cell clumps remain.

5. Add 4 ml Buffer P2, mix thoroughly by vigorously inverting the sealed tube 4–6 times, and incubate at room temperature (15–25°C) for 5 min. Total volume is now 8ml.

Do not vortex, as this will result in shearing of genomic DNA. The lysate should appear viscous. Do not allow the lysis reaction to proceed for more than 5 min. After use, the bottle containing Buffer P2 should be closed immediately to avoid acidification from CO₂ in the air.

If LyseBlue has been added to Buffer P1, the cell suspension will turn blue after addition of Buffer P2. Mixing should result in a homogeneously colored suspension. If the suspension contains localized colorless regions or if brownish cell clumps are still visible, continue mixing the solution until a homogeneously colored suspension is achieved.

6. Add 4 ml of chilled Buffer P3, mix immediately and thoroughly by vigorously inverting 4–6 times, and incubate on ice for 15 min. Total volume is now 12ml.

Precipitation is enhanced by using chilled Buffer P3 and incubating on ice. After addition of Buffer P3, a fluffy white material forms and the lysate becomes less viscous. The precipitated material contains genomic DNA, proteins, cell debris, and KDS. The lysate should be mixed thoroughly to ensure even potassium dodecyl sulfate precipitation. If the mixture still appears viscous, more mixing is required to completely neutralize the solution. If LyseBlue reagent has been used, the suspension should be mixed until all trace of blue has gone and the suspension is colorless. A homogeneous colorless suspension indicates that the SDS has been effectively precipitated.

7. Centrifuge 12 ml volume at $\geq 20,000 \times g$ for 30min at 4°C. Remove supernatant containing plasmid DNA promptly. Use high-speed centrifuge (in BSL3 or equivalent). If BSL3 centrifuge is utilized, all steps there after must take place within the BSL3.

Before loading the centrifuge, the sample should be mixed again. Centrifugation should be performed in non-glass tubes (e.g., polypropylene). After centrifugation the supernatant should be clear.

8. Centrifuge the supernatant again at $\geq 20,000 \times g$ for 15 min at 4°C. Remove supernatant containing plasmid DNA promptly.

This second centrifugation step should be carried out to avoid applying suspended or particulate material to the QIAGEN-tip. Suspended material (causing the sample to appear turbid) can clog the QIAGEN-tip and reduce or eliminate gravity flow.

9. Equilibrate a QIAGEN-tip 100 by applying 4 ml Buffer QBT, and allow the column to empty by gravity flow. Prepare Qiagen-Tip 100 during centrifugation by adding 4ml of Buffer QBT.

Flow of buffer will begin automatically by reduction in surface tension due to the presence of detergent in the equilibration buffer. Allow the QIAGEN-tip to drain completely. QIAGEN-tips can be left unattended, since the flow of buffer will stop when the meniscus reaches the upper frit in the column.

10. Apply the supernatant from step 8 to the QIAGEN-tip immediately after centrifugation and allow it to enter the resin by gravity flow.

The supernatant should be loaded onto the QIAGEN-tip promptly. If it is left too long and becomes cloudy due to further precipitation of protein, it must be centrifuged again or filtered before loading to prevent clogging of the QIAGEN-tip.

11. Wash the QIAGEN-tip twice with 10 ml Buffer QC.

Allow Buffer QC to move through the QIAGEN-tip by gravity flow. The first wash is sufficient to remove contaminants in the majority of plasmid DNA preparations. The second wash is especially necessary when large culture volumes or bacterial strains producing large amounts of carbohydrates are used.

12. Elute DNA with 5 ml Buffer QF into a centrifuge tube.

Collect the eluate in a 15 ml or 50 ml tube (not supplied). Use of polycarbonate centrifuge tubes is not recommended as polycarbonate is not resistant to the alcohol used in subsequent steps. For constructs larger than 45–50 kb, prewarming the elution buffer to 65°C may help to increase yield.

Optional: If you wish to stop the protocol and continue later, store the eluate at 4°C. Storage periods longer than overnight are not recommended.

13. Precipitate DNA by adding 3.5 ml (0.7 volumes) room-temperature isopropanol to the eluted DNA. Mix and centrifuge immediately at $\geq 15,000 \times g$ for 30 min at 4°C. Carefully decant the supernatant.

All solutions should be at room temperature to minimize salt precipitation, although centrifugation is carried out at 4°C to prevent overheating of the sample. Alternatively, disposable conical bottom centrifuge tubes can be used for centrifugation at $5000 \times g$ for 60 min at 4°C. Isopropanol pellets have a glassy appearance and may be more difficult to see than the fluffy, salt-containing pellets that result from ethanol precipitation. Marking the outside of the tube before centrifugation allows the pellet to be more easily located. Isopropanol pellets are also more loosely attached to the side of the tube, and care should be taken when removing the supernatant.

14. Wash DNA pellet with 2 ml of room-temperature 70% ethanol, and centrifuge at $\geq 15,000 \times g$ for 10 min. Carefully decant the supernatant without disturbing the pellet.

Alternatively, disposable conical-bottom centrifuge tubes can be used for centrifugation at $5000 \times g$ for 60 min at 4°C. The 70% ethanol removes precipitated salt and replaces isopropanol with the more volatile ethanol, making the DNA easier to redissolve.

15. Air-dry the pellet for 5–10min, and redissolve the DNA in a suitable volume of buffer (e.g., TE buffer, pH 8.0, or 10 mM Tris-Cl, pH 8.5). Add approx. 100µl of TE Buffer to dissolve the plasmid DNA.

Redissolve the DNA pellet by rinsing the walls to recover the DNA, especially if glass tubes have been used. Pipetting the DNA up and down to promote resuspension may cause shearing and should be avoided. Overdrying the pellet will make the DNA difficult to redissolve. DNA dissolves best under slightly alkaline conditions; it does not easily dissolve in acidic buffers.

16. Plate the Plasmid onto a Blood Agar Plate and incubate for 72hrs. Plate may be checked every 24hrs for growth. If no growth is observed after 72hrs, the plasmid can be removed from the BSL3.

Quantification

1. Determine the concentration of plasmid DNA recovered using the Thermo Scientific NanoDrop instrument (or other spectrophotometer). Further dilutions can be made with TE Buffer to achieve target concentration of 40mM.
2. Additional dilutions should be made according to the *Vibrio parahaemolyticus* and *Vibrio vulnificus* enumeration and detection through MPN and real-time PCR assay protocol.
3. Plasmid dilutions must be run on a PCR Detection System to verify concentration, and ensure the amplification falls within its expected Ct range.
4. 10µl aliquots of 1:100 dilution will serve as freezer stock and be stored at -15°C or below. Working stocks can be prepared by adding 990µl molecular grade water to the freezer stock, and should be stored at 2-8°C. On day of use, dilute working stock an additional 1:100. The final concentration of the final product is 1:100,000.

Validation Data for MPN Real-time PCR for Total and Pathogenic *Vibrio parahaemolyticus*

Name of Method Submitter: Gina Olson, Washington State Department of Health

Specific purpose or intent of the method for use in the NSSP:

Requesting adoption of this method as an approved method for *Vibrio* enumeration, both *Vibrio parahaemolyticus* (Vp) and *Vibrio vulnificus* (Vv) in oysters. The method may be used in the following applications: PHP validation and verification of product and in management of growing areas through environmental testing and surveillance in order to re-open closed growing areas. This method once approved would provide a high-throughput alternative to the current approved MPN real-time PCR method. In addition, this method would be the only approved MPN real-time PCR method to test for total Vp, pathogenic Vp, and Vv in a single assay.

Validation Criteria Data:

All oyster samples used in this validation were collected from different harvest locations and/or different harvest dates in Washington State. All samples were processed using APHA guidelines for the cleaning and shucking of shellstock. Samples were confirmed negative for the target organism of Vp through the FDA BAM culture-based method and through pcr prior to spiking. Spiking levels were determined by spread plating dilutions onto PCA w/2% NaCl in duplicate and averaging the counts.

Vp strain WA4647 was used to spike all samples for all validation criteria. This strain is positive for *tlh*, *tdh*, and *trh*. All data generated for all three targets was identical and has been presented in a single MPN in all validation criteria tables and data calculations.

The validation data for Vp and Vv is presented separately for clarity and ease in reviewing the data, but this is a single assay and all elements were present during the validation of all organisms.

Assay Design

DNA Isolation: Roche MagnaPure 96 using Roche DNA/Viral Nucleic Acid Small Volume Kit

Real-time PCR Instrument: Applied Biosystems QuantStudio Dx (384-well format)

Mastermix: Life Technologies TaqMan Environmental Master Mix 2.0

Real-Time PCR targets: 2 multiplex reactions

- Multiplex 1: Total Vp (*tlh*), Vv (*vvhA*), internal control (IC)
- Multiplex 2 (Vp pathogenicity markers): *tdh*, *trh*, *orf8*

Real-Time PCR parameters:

Denaturation: 95°C for 10 mins
Annealing: 95°C for 15 secs
Extension: 59°C for 1 min
Cycles: 40

1. Accuracy/Trueness & Measurement Uncertainty

Accuracy/Trueness

Purpose/Method

Accuracy/Trueness measures the closeness of agreement between the test results (MPNs) and the accepted reference values (CFUs). This was done by analyzing twenty oyster samples over a range of concentrations (low to high) to determine the MPN. The MPN and CFU data set was converted into logs. The average MPN in logs was divided by the average plate count in logs. This provides an estimate (in percent) of the accuracy/trueness of the method.

Results

The average of the plate count CFUs was 2.88 log. The average of MPNs was 3.17 log. Accuracy/Trueness was found to be 109.94%. Results can be found below in Table 1.

Measurement Uncertainty

Purpose/Method

Measurement uncertainty expresses the range of values around the measured result within which the true value is expected to lie. To determine this parameter, twenty oyster samples spiked with a range of concentrations were analyzed. The MPN and CFU data set was converted into logs and the MPN result was subtracted from the CFU result for each sample. A 95% confidence interval was calculated from the difference. This confidence interval represents the measurement uncertainty of the methods.

Results

The measurement uncertainty was determined via 95% CI (0.23, 0.50), resulting in a measurement uncertainty of 0.27. Results can be found below in Table 1.

Table 1. Data for determination of Accuracy/Trueness and Measurement Uncertainty

Sample	Plate Count, log(CFUs)	MPN, log(MPN/g)
1	0.89	0.62
2	0.77	1.36
3	0.85	0.96
4	2.85	2.62
5	2.32	3.36
6	2.32	3.17
7	4.18	5.04
8	4.18	4.66
9	0.36	0.63

10	4.36	4.66
11	2.04	2.62
12	2.45	2.36
13	3.52	3.36
14	4.69	5.04
15	4.69	5.04
16	1.66	1.62
17	2.57	2.96
18	3.60	3.59
19	4.69	5.04
20	4.69	4.66

2. Ruggedness

Purpose/Method

The amount of analyte recovered should be consistent between different lots of media/reagents. Ruggedness tests the impact of different lots used to process samples on the final result. This was done by testing ten oyster samples spiked at a range of concentrations in duplicate. The first replicate was performed using "Lot 1" media/reagents and the second was performed using "Lot 2." To determine if the method was sufficiently rugged to withstand the types of changes anticipated to occur in routine use, a two-sided t-test was utilized on log-transformed data with a significance level (α) of 0.05. There should be no significant difference between Lot 1 and Lot 2 samples.

Results

Using data from Table 2, there was no significant difference ($p=0.64$) between different lots of media and reagents.

Table 2. Data for Determination of Ruggedness

Sample	Replicate 1, log(MPN/g)	Replicate 2, log(MPN/g)
1	0.62	0.96
2	1.36	1.36
3	0.96	0.96
4	2.96	2.62

5	3.36	3.63
6	2.96	3.18
7	5.04	5.34
8	5.34	4.66
9	0.62	1.62
10	5.04	4.66

3. Precision & Recovery

Precision

Purpose/Method

The difference between the methods results (MPNs) and the reference values (CFUs) should be consistent between different samples and also when detecting varying concentrations of measurand. The precision of the method tests the consistency of the difference between the CFU's found on plates and the MPN values. This was done by testing ten oyster samples at low, medium, and high concentrations in duplicate to determine the MPN. The MPN and CFU data sets were converted into log values. Each MPN was compared to its associate CFU value. A nested ANOVA was then performed, with variance components being the sample, and concentrations within the samples (low, medium, and high), and then the error. The ANOVA component of interest was that comparing the concentrations within the samples to the determinations (or error).

Results

The difference between CFUs and MPNs can be found below in Table 3. The data shows that there are no significant differences between the concentrations in samples and the determinations within concentrations via a nested ANOVA ($p=0.08$). Additionally, the variance of the method does not exceed the known variance of a 3-tube MPN ($p=0.21$).

Table 3: Data for determining the Precision

Sample	Concentration	Difference	log(CFUs)	log(MPNs)
1	low	-0.72	-1.17	-0.44
1	low	-1.13	-1.17	-0.04
1	med	-1.40	1.96	3.36
1	med	-1.00	1.96	2.96
1	high	0.13	4.69	4.56

1	high	-0.65	4.69	5.34
2	low	-0.68	-0.51	0.18
2	low	0.24	-0.51	-0.75
2	med	0.08	2.45	2.36
2	med	-0.42	2.45	2.87
2	high	-0.35	4.69	5.04
2	high	0.03	4.69	4.66
3	low	-0.18	-0.92	-0.74
3	low	-0.18	-0.92	-0.74
3	med	-0.45	2.52	2.96
3	med	-0.11	2.52	2.62
3	high	-0.35	4.69	5.04
3	high	-0.35	4.69	5.04
4	low	-0.08	-0.21	-0.13
4	low	-0.57	-0.21	0.36
4	med	-0.13	2.49	2.62
4	med	-0.13	2.49	2.62
4	high	-0.35	4.69	5.04
4	high	0.03	4.69	4.66
5	low	0.07	-0.15	-0.21
5	low	0.30	-0.15	-0.44
5	med	0.13	2.49	2.36
5	med	0.13	2.49	2.36
5	high	-0.35	4.69	5.04
5	high	0.03	4.69	4.66
6	low	-0.24	-0.28	-0.04

6	low	0.17	-0.28	-0.44
6	med	-0.30	2.66	2.96
6	med	0.30	2.66	2.36
6	high	-0.35	4.69	5.04
6	high	-0.35	4.69	5.04
7	low	0.34	-0.41	-0.74
7	low	0.34	-0.41	-0.74
7	med	-0.40	2.57	2.96
7	med	-0.40	2.57	2.96
7	high	0.03	4.69	4.66
7	high	-0.65	4.69	5.34
8	low	-0.12	-0.57	-0.44
8	low	-0.44	-0.57	-0.13
8	med	-0.02	2.60	2.62
8	med	-0.02	2.60	2.62
8	high	0.03	4.69	4.66
8	high	0.03	4.69	4.66
9	low	0.12	-0.33	-0.44
9	low	-0.20	-0.33	-0.13
9	med	-0.12	2.52	2.62
9	med	-0.45	2.52	2.96
9	high	-0.35	4.69	5.04
9	high	-0.35	4.69	5.04
10	low	0.03	-0.48	-0.51
10	low	0.03	-0.48	-0.51
10	med	-0.29	2.67	2.96

10	med	-0.96	2.67	3.63
10	high	0.03	4.69	4.66
10	high	0.03	4.69	4.66

Recovery

Purpose/Method

The amount of analyte recovered should be consistent both between different samples and also when detecting varying concentrations of measurand. The recovery of the method tests the consistency of the analyte recovered via MPNs as compared to the CFUs found on plates. This was done by testing ten oyster samples at low, medium, and high concentrations in duplicate to determine the MPN. The MPN and CFU data set was converted to logs. Each duplicated MPN was averaged and then compared to its associate CFU, in logs. A single-factor ANOVA was then used to compare the recovery at the three concentrations.

Results

The difference between CFUs and MPNs can be found below in Table 4. The recovery across all samples and concentrations was found to be 109.71%. There was not found to be significant differences in the recovery at the various concentrations ($p=0.56$).

Table 4: Data for the determination of Recovery

Sample	Concentration	Ave log(CFUs) per Conc.	Ave log(MPN) per Conc.
1	low	-1.17	-0.24
	med	1.96	3.16
	high	4.69	4.95
2	low	-0.51	-0.28
	med	2.45	2.62
	high	4.69	4.85
3	low	-0.92	-0.75
	med	2.52	2.79
	high	4.69	5.04
4	low	-0.21	0.12
	med	2.49	2.62
	high	4.69	4.85
5	low	-0.15	-0.33
	med	2.49	2.36
	high	4.69	4.85
6	low	-0.28	-0.24
	med	2.66	2.66
	high	4.69	5.04

7	low	-0.41	-0.75
	med	2.57	2.96
	high	4.69	5.00
8	low	-0.57	-0.29
	med	2.6	2.62
	high	4.69	4.66
9	low	-0.33	-0.29
	med	2.52	2.79
	high	4.69	5.04
10	low	-0.48	-0.51
	med	2.67	3.3
	high	4.69	4.66

4. Specificity

Purpose/Method

The method should only detect the analyte of interest, even in the presence of interfering organisms. Specificity refers to the ability of the method to measure only the target organism. One matrix sample was divided into three aliquots. One aliquot was spiked with a low but determinable level of *Vibrio parahaemolyticus* (*Vp*). The other two aliquots were spiked with the same level of *Vp* as the first, but were also spiked with a high level of potential interfering organisms. One aliquot received a high level of *Vibrio vulnificus* (*Vv*) and the other received *Vibrio alginolyticus* (*Va*). Five replicates were performed. Each of the replicates was analyzed by taking the average log MPN and calculating the Specificity Index (SI). A paired t-test was used to determine if the average specificity index obtained from the five replicates differed from 1 (significance level = 0.05).

Results

Using the data from Table 5, the average specificity index was 0.88 when in the presence of *Vv* and 0.98 in the presence of *Va*. These values are not significantly different than 1 ($p=0.12$ for *Vv*, $p=0.69$ for *Va*).

Table 5. Data for Determination of Specificity

Replicate	<i>Vp</i> only, log(MPN/g)	<i>Vp</i> + <i>Vv</i> , log(MPN/g)	<i>Vp</i> + <i>Va</i> , log(MPN/g)
1	1.96	2.36	2.36
2	1.62	1.58	2.36
3	1.62	1.96	1.96
4	2.36	2.36	1.96
5	2.17	2.96	1.62

5. Linear Range, Limit of Detection & Limit of Quantification/Sensitivity

Linear Range

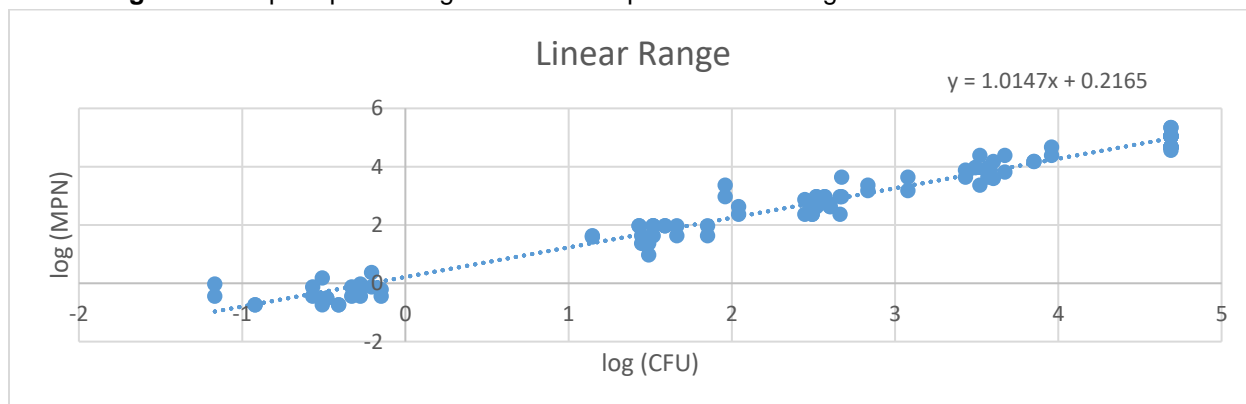
Purpose/Method

The MPN value found should directly correlate to the concentration of analyte within the sample, within the working range of the method. Thus, as the concentration increases, the MPN value should also increase in a linear fashion. Ten Oyster samples were tested at 5 concentration levels, in duplicate. Each MPN was compared to its associate CFU, found by plate count. The relationship between the log(MPN) and log(CFU) was then found by obtaining the correlation coefficient by performing a linear regression with log(CFU) as the independent variable and log(MPN) as the dependent variable.

Results

The relationship between the MPNs and CFUs can be seen in Figure 1 below. The relationship between MPNs and CFUs was found to be linear, with a Pearson's r of 0.99. The working range used was of concentrations ranging from 10^{-1} to 10^4 cells/gram.

Figure 1: Graph representing the relationship between the log values of CFUs and MPNS



Limit of Detection

Purpose/Method

The method should be capable of detecting as little as 1 cell/gram of sample, or 0 cells/gram, in log form. Therefore, it must be determined whether the method can detect one cell per gram of sample. The log(MPN) was compared to the log(CFU) of ten oyster samples, spiked at five varying concentrations, in duplicate. This was done by performing a regression analysis on the data and calculating the Limit of Detection by taking the antilog of the intercept. The independent variable was set as log(CFU) and the dependent variable was set as log(MPN).

Results

The Limit of Detection was found to be 1.65 cells. The overall regression standard error, the

95.0% confidence interval was found to be 0.67, which encompasses the Limit of Detection. The 99.99% confidence interval of the intercept was found to be 0.23. These both contain the intercept of 0.22 within the interval.

Table 7: Data for determination of the Limit of Detection

Sample	Concentration	Log(CFU)	Log(MPN)
Sample 1, Rep 1	10⁻¹	-1.17	-0.45
	10¹	2.05	2.63
	10²	2.96	3.36
	10³	3.96	4.66
	10⁴	4.69	4.66
Sample 1, Rep 2	10⁻¹	-1.17	-0.04
	10¹	2.05	2.36
	10²	2.96	2.96
	10³	3.96	4.38
	10⁴	4.69	5.34
Sample 2, Rep 1	10⁻¹	-0.51	0.17
	10¹	1.44	1.36
	10²	2.44	2.36
	10³	2.83	3.17
	10⁴	4.69	5.04

Sample 2, Rep 2	10⁻¹	-0.51	-0.75
	10¹	1.44	1.63
	10²	2.44	2.87
	10³	2.83	3.36
	10⁴	4.69	4.66
Sample 3, Rep 1	10⁻¹	-0.93	-0.75
	10¹	1.13	1.58
	10²	2.51	2.96
	10³	3.51	3.36
	10⁴	4.69	5.04
Sample 3, Rep 2	10⁻¹	-0.93	-0.75
	10¹	1.13	1.63
	10²	2.51	2.63
	10³	3.51	4.38
	10⁴	4.69	5.04
Sample 4, Rep 1	10⁻¹	-0.21	-0.13
	10¹	1.48	0.96
	10²	2.48	2.63

	10³	3.07	3.63
	10⁴	4.69	5.04
Sample 4, Rep 2	10⁻¹	-0.21	0.36
	10¹	1.48	1.36
	10²	2.48	2.63
	10³	3.07	3.17
	10⁴	4.69	4.66
Sample 5, Rep 1	10⁻¹	-0.15	-0.21
	10¹	1.85	1.96
	10²	2.49	2.36
	10³	3.49	3.97
	10⁴	4.69	5.04
Sample 5, Rep 2	10⁻¹	-0.15	-0.45
	10¹	1.85	1.63
	10²	2.49	2.36
	10³	3.49	3.97
	10⁴	4.69	4.66
Sample 6, Rep 1	10⁻¹	-0.28	-0.04

	10¹	1.66	1.63
	10²	2.66	2.96
	10³	3.85	4.17
	10⁴	4.69	5.04
Sample 6, Rep 2	10⁻¹	-0.28	-0.45
	10¹	1.66	1.96
	10²	2.66	2.36
	10³	3.85	4.17
	10⁴	4.69	5.04
Sample 7, Rep 1	10⁻¹	-0.41	-0.75
	10¹	1.59	1.96
	10²	2.57	2.96
	10³	3.57	3.97
	10⁴	4.69	4.66
Sample 7, Rep 2	10⁻¹	-0.41	-0.75
	10¹	1.59	1.96
	10²	2.57	2.96
	10³	3.57	3.63

	10⁴	4.69	5.34
Sample 8, Rep 1	10⁻¹	-0.57	-0.45
	10¹	1.43	1.96
	10²	2.60	2.63
	10³	3.60	3.59
	10⁴	4.69	4.66
Sample 8, Rep 2	10⁻¹	-0.57	-0.13
	10¹	1.43	1.96
	10²	2.60	2.63
	10³	3.60	4.17
	10⁴	4.69	4.66
Sample 9, Rep 1	10⁻¹	-0.33	-0.45
	10¹	1.51	1.96
	10²	2.51	2.63
	10³	3.43	3.63
	10⁴	4.69	5.04
Sample 9, Rep 2	10⁻¹	-0.33	-0.13
	10¹	1.51	1.63

	10²	2.51	2.96
	10³	3.43	3.87
	10⁴	4.69	5.04
Sample 10, Rep 1	10⁻¹	-0.49	-0.52
	10¹	1.51	1.96
	10²	2.67	2.96
	10³	3.67	4.38
	10⁴	4.69	4.66
Sample 10, Rep 2	10⁻¹	-0.49	-0.52
	10¹	1.51	1.96
	10²	2.67	3.63
	10³	3.67	3.80
	10⁴	4.69	4.66

Limit of Quantification/Sensitivity

Purpose/Method

The quantifiable limit of the method is bounded by the values defined by a 3-tube MPN. In the case that the Limit of Detection is not significantly different than 1 cell/gram, than the Limit of Quantification can be extrapolated using the FDA BAM MPN Calculator.

Results

As the method starts with a low dilution of 1 gram of sample per tube, use of a 3-tube MPN and corresponding dilution ratios will result in the Limit of Quantification/Sensitivity for the method being 0.36 MPN/gram.

Inclusivity

Purpose

To assess the ability of the method to detect a wide range of target strains in various oyster tissues.

Method

Vibrio parahaemolyticus (*Vp*) strains were grown in APW for 18-24 hours. DNA was extracted using the MagNAPure 96 instrument and PCR performed using the QuantStudio Dx Real-Time PCR System. The strains that make up the inclusivity panel were obtained from the Center for Disease Control and Prevention (CDC), National Oceanic and Atmospheric Administration (NOAA), American Type Culture Collection (ATCC), or Washington State Public Health Laboratory (WAPHL). The primers and probes utilized in this method have been tested against DNA extracts from the isolates listed in Table 1 below.

Confirmation was performed through a secondary assay. Since *tlh* is a species specific marker, a positive *Vp* identification through biochemicals or by ATCC paperwork was used as confirmation. For *tdh* some isolates were confirmed by NOAA and others were confirmed using *tdh* primers from Nordstrom et al. 2007. The *trh* marker was more challenging due to 2 variations of the *trh* gene. All isolates were confirmed using the FDA BAM *trh* primer set and anything with discrepant results between our assay and the FDA assay we confirmed with a biochemical urease test. The ability of *Vp* to hydrolyze urea has been shown to be indicative of the presence of the *trh* gene (Lida et al paper 1997). The *ORF8* pandemic marker was confirmed using the primer set from Myers et al. 2003.

Results

Primer / Probe Sensitivity

Sensitivity= (# of true positives/ (# of true positives + # of false negatives))

tlh sensitivity = 73/73 = **100% *tlh* sensitivity**

tdh sensitivity = 33/33 = **100% *tdh* sensitivity**

trh sensitivity = 33/33 = **100% *trh* sensitivity**

ORF8 sensitivity = 24/24 = **100% *ORF8* sensitivity**

The primers and probes utilized in this method for their respective target demonstrates 100% inclusivity. See Table 1 (Inclusivity Panel) for the breakdown of strains, source, targets tested, and its corresponding result.

Table 1. Inclusivity panel with results by target. Targets not tested or confirmed are blocked out.

Strain	Source	tlh	tdh	trh	orf8	vvha
F5828	CDC	+	+		+	
F5835	CDC	+	+		+	
F5847	CDC	+	+		+	
F6820	CDC	+	+		+	
F7630	CDC	+	+		+	
F7635	CDC	+	+		+	
F7636	CDC	+	+		+	
F7680	CDC	+	+		+	
F8701	CDC	+	+		+	
F8949	CDC	+	+		+	
F9083	CDC	+	+		+	
K0071	CDC	+	+		+	
K0456	CDC	+	-	+	-	
17803	ATCC	+	-	+		
27519	ATCC	+	-			
27969	ATCC	+				
33844	ATCC	+	+			
33845	ATCC	+	+			
33846	ATCC	+	+			
33847	ATCC	+	+			
35117	ATCC	+	-			
35118	ATCC	+	+			
43996	ATCC	+	+			
49398	ATCC	+	-			
BAA-238	ATCC	+	+		+	
BAA-239	ATCC	+	+		+	
BAA-240	ATCC	+	+		+	
BAA-241	ATCC	+	+		+	
BAA-242	ATCC	+	+		+	
NWF 261	NOAA - NWFSC	+	-	-	-	
NWF 512	NOAA - NWFSC	+	-	-	-	
NWF 586	NOAA - NWFSC	+	+	-	+	
NWF 605	NOAA - NWFSC	+	+	-	+	
NWF 609	NOAA - NWFSC	+	+	-	+	
NWF 735	NOAA - NWFSC	+	+		+	
NWF 782	NOAA - NWFSC	+	+	-	+	

NWF 797	NOAA - NWFSC	+	-	-		
NWF 800	NOAA - NWFSC	+	-			
NWF 805	NOAA - NWFSC	+	-			
NWF 843	NOAA - NWFSC	+	-			
NWF 846	NOAA - NWFSC	+	+	+		
NWF 864	NOAA - NWFSC	+	+		+	
NWF 930	NOAA - NWFSC	+	+	-	+	
5412	WA PHL	+		+		
5419	WA PHL	+		+		
5423	WA PHL	+	+	+		
5424	WA PHL	+		+		
5425	WA PHL	+	+	+		
5426	WA PHL	+		+		
5429	WA PHL	+		+		
5430	WA PHL	+		+		
5434	WA PHL	+		+		
5436	WA PHL	+		+		
5437	WA PHL	+		+		
5442	WA PHL	+		+		
5444	WA PHL	+		+		
5454	WA PHL	+		+		
5456	WA PHL	+		+		
5463	WA PHL	+		+		
5468	WA PHL	+		+		
5469	WA PHL	+		+		
5470	WA PHL	+		+		
5471	WA PHL	+		+		
5473	WA PHL	+		+		
5474	WA PHL	+		+		
5475	WA PHL	+		+		
5487	WA PHL	+		+		
5488	WA PHL	+		+		
5492	WA PHL	+		+		
5501	WA PHL	+		+		
5508	WA PHL	+		+		
5518	WA PHL	+		+		
5519	WA PHL	+		+		
06-2410	06-2410 (CDC)					+

06-2450	06-2450 (CDC)					+
07-2405	07-2405 (CDC)					+
08-2468	08-2468 (CDC)					+
08-2470	08-2470 (CDC)					+
08-2472	08-2472 (CDC)					+
08-2485	08-2485 (CDC)					+
1831-81	1831-81 (CDC)					+
2009V-1002	2009V-1002 (CDC)					+
2009V-1055	2009V-1055 (CDC)					+
2010V-1021	2010V-1021 (CDC)					+
209V-1035	209V-1035 (CDC)					+
2431-04	2431-04 (CDC)					+
2473-85	2473-85 (CDC)					+
2492-88	2492-88 (CDC)					+
2809-78	2809-78 (CDC)					+
430-79	430-79 (CDC)					+
AM38622	AM38622 (CDC)					+
AM38623	AM38623 (CDC)					+
27562	27562					+
29307	29307					+
Total Confirmed Isolated		73	33	33	24	21

Exclusivity

Purpose

To demonstrate the ability of the method to distinguish the targeted analyte from other potentially cross-reactive non-target strains that could possibly contaminate shellfish.

Method

All organisms were inoculated into APW and incubated for 18-24 hours. DNA was extracted using the MagNAPure 96 instrument and PCR performed using the QuantStudio Dx Real-Time PCR System. All strains were obtained from the Center for Disease Control and Prevention (CDC) and American Type Culture Collection (ATCC).

Results

Primer / Probe Specificity

Specificity = (# of true negative/ (# of true negatives + # of false positives))

tlh specificity= 49/49 = **100% *tlh* Specificity**

tdh specificity = 49/49 = **100% *tdh* Specificity**

trh specificity = 49/50 = **98% *trh* Specificity**

ORF8 specificity = 49/49 = **100% *ORF8* Specificity**

The *tlh*, *tdh*, and *ORF8* primers and probes utilized in this method demonstrate 100% exclusivity. The *trh* primers and probe demonstrate a 98% specificity (see Known Limitations below). See Table 2 (Exclusivity Panel) for the breakdown of strains, source, targets tested, and its corresponding result.

Known limitations and interferences

Vibrio alginolyticus possesses a *trh* gene with 98% homology to the *trh* gene in *Vibrio parahaemolyticus*. Most probable number (MPN) values for *trh* should be reported only if *tlh* (*V. parahaemolyticus* specific gene) is present in the corresponding tube. Tubes only positive for *trh* should not be accounted for when generating the MPN value.

This assay utilizes the Taqman Environmental Mastermix 2.0, which is specifically formulated to detect bacterial pathogens with greater specificity and sensitivity. There are no additional known limitations when using the Taqman Environmental Mastermix 2.0.

Table 2. Exclusivity panel with results by target. Targets not tested or confirmed are blocked out.

Organism	Strain	<i>tlh</i>	<i>tdh</i>	<i>trh</i>	<i>orf8</i>	<i>vvha</i>
<i>E. coli</i>	ATCC 25922	-	-	-	-	-
<i>G. hollisae</i>	ATCC 33564	-	-	-	-	-
<i>K. pneumoniae</i>	ATCC 33495	-	-	-	-	-
<i>P. aeruginosa</i>	ATCC 33495	-	-	-	-	-
<i>S. aureus</i>	ATCC 10145	-	-	-	-	-
<i>S. sonnei</i>	ATCC 25925	-	-	-	-	-
<i>S. typhimurium</i>	ATCC 9290	-	-	-	-	-
<i>V. aestuarians</i>	ATCC 35048	-	-	-	-	-
<i>V. alginolyticus</i>	ATCC 17749	-	-	-	-	-
<i>V. alginolyticus</i>	S14-048 (Environmental- WA PHL)	-	-	+	-	-
<i>V. alginosus</i>	ATCC 14390	-	-	-	-	-
<i>V. campbellii</i>	ATCC 25920	-	-	-	-	-
<i>V. cholerae</i>	ATCC 39050	-	-	-	-	-
<i>V. cincinnatiensis</i>	ATCC 35912	-	-	-	-	-
<i>V. furnissii</i>	ATCC 33813	-	-	-	-	-
<i>V. marinagilis</i>	ATCC 14398	-	-	-	-	-

<i>V. marinofulvus</i>	ATCC 14395	-	-	-	-	-
<i>V. marinovulgaris</i>	ATCC 14394	-	-	-	-	-
<i>V. metschnikovii</i>	ATCC 700040	-	-	-	-	-
<i>V. mimicus</i>	ATCC 33653	-	-	-	-	-
<i>V. natriegens</i>	ATCC 14048	-	-	-	-	-
<i>V. nereis</i>	ATCC 25917	-	-	-	-	-
<i>V. nigripulchritudo</i>	ATCC 27043	-	-	-	-	-
<i>V. ponticus</i>	ATCC 14391	-	-	-	-	-
<i>V. proteolyticus</i>	ATCC 15338	-	-	-	-	-
<i>V. spledidus</i>	ATCC 33789	-	-	-	-	-
<i>V. tubiashii</i>	ATCC 19106	-	-	-	-	-
<i>V. vulnificus</i>	06-2410 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	06-2450 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	07-2405 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	08-2468 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	08-2470 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	08-2472 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	08-2485 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	1831-81 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	2009V-1002 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	2009V-1055 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	2010V-1021 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	209V-1035 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	2431-04 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	2473-85 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	2492-88 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	2809-78 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	430-79 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	AM38622 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	AM38623 (CDC)	-	-	-	-	
<i>V. vulnificus</i>	27562	-	-	-	-	
<i>V. vulnificus</i>	29307	-	-	-	-	
<i>A. trota</i>	2013V-1197 (CDC)	-	-	-	-	-
<i>A. veronii</i>	N/A (CDC)	-	-	-	-	-
Total Strains		49				

Validation Data for MPN Real-time PCR for *Vibrio vulnificus*

Name of Method Submitter: Gina Olson, Washington State Department of Health

Specific purpose or intent of the method for use in the NSSP:

Requesting adoption of this method as an approved method for *Vibrio* enumeration, both *Vibrio parahaemolyticus* (Vp) and *Vibrio vulnificus* (Vv) in oysters. The method may be used in the following applications: PHP validation and verification of product and in management of growing areas through environmental testing and surveillance in order to re-open closed growing areas. This method once approved would provide a high-throughput alternative to the current approved MPN real-time PCR method. In addition, this method would be the only approved MPN real-time PCR method to test for total Vp, pathogenic Vp, and Vv in a single assay.

Validation Criteria Data:

All oyster samples used in this validation were collected from different harvest locations and/or different harvest dates in Washington State. All samples were processed using APHA guidelines for the cleaning and shucking of shellstock. Samples were confirmed negative for the target organism of Vv through the FDA BAM culture-based method and through pcr prior to spiking. Spiking levels were determined using a 5-tube MPN dilution series in APW in duplicate (averaging the 2 values).

Vv strain ATCC 29307 was used to spike all samples for all validation criteria. This strain is positive for *vvhA*.

The validation data for Vp and Vv is presented separately for clarity and ease in reviewing the data, but this is a single assay and all elements were present during the validation of all organisms.

Assay Design

DNA Isolation: Roche MagnaPure 96 using Roche DNA/Viral Nucleic Acid Small Volume Kit

Real-time PCR Instrument: Applied Biosystems QuantStudio Dx (384-well format)

Mastermix: Life Technologies TaqMan Environmental Master Mix 2.0

Real-Time PCR targets: 2 multiplex reactions

- Multiplex 1: Total Vp (*tlh*), Vv (*vvhA*), internal control (IC)
- Multiplex 2 (Vp pathogenicity markers): *tdh*, *trh*, *orf8*

Real-Time PCR parameters:

Denaturation: 95°C for 10 mins

Annealing: 95°C for 15 secs

Extension: 59°C for 1 min

Cycles: 40

1. Accuracy/Trueness & Measurement Uncertainty

Accuracy/Trueness

Purpose/Method

Accuracy/Trueness measures the closeness of agreement between the test results (MPNs) and the reference results (spiked MPNs without matrix). This was done by analyzing twenty oyster samples over a range of concentrations (low to high) to determine the MPN. The MPN and reference data set was converted into logs. The average MPN in logs was divided by the average reference value in logs. This provides an estimate in percent of the accuracy/trueness of the method.

Results

The average of the reference values was 2.20 log. The average of MPNs was 2.15 log. Accuracy/Trueness was found to be 97.69%. Results can be found below in Table 1.

Measurement Uncertainty

Purpose/Method

Measurement uncertainty expresses the range of values around the measured result within which the true value is expected to lie. To determine this parameter, twenty oyster samples spiked with a range of concentrations were analyzed. Each MPN and reference value was converted into logs and the MPN result was subtracted from the reference result for each sample. A 95% confidence interval was calculated from the difference. This confidence interval represents the measurement uncertainty of the methods.

Results

The measurement uncertainty was determined via 95% CI (0.16, 0.30), resulting in a measurement uncertainty of 0.14. Results can be found below in Table 1.

Table 1. Data for determination of Accuracy/Trueness and Measurement Uncertainty

Sample	Reference MPN, log(MPN/g)	MPN, log(MPN/g)
1	0.11	-0.44
2	0.11	-0.04
3	0.11	0.36
4	2.11	1.96
5	2.11	2.36
6	2.20	1.87

7	2.20	2.36
8	4.08	3.97
9	4.08	4.38
10	4.08	3.88
11	0.23	-0.04
12	1.18	1.18
13	2.30	1.96
14	3.15	3.36
15	4.30	4.38
16	0.49	0.36
17	1.32	0.96
18	2.08	2.62
19	3.45	3.36
20	4.30	4.18

2. Ruggedness

Purpose/Method

The amount of analyte recovered should be consistent between different lots of media/reagents. Ruggedness tests the impact of different lots used to process samples on the final result. This was done by testing ten oyster samples spiked at a range of concentrations in duplicate. One replicate was performed using "Lot 1" media/reagents and one replicate was performed using "Lot 2." To determine if the method was sufficiently rugged to withstand the types of changes anticipated to occur in routine use, a two-sided t-test was utilized on log-transformed data with a significance level (α) of 0.05. There should be no significant difference between Lot 1 and Lot 2 samples.

Results

Using data from Table 2, there was no significant difference ($p=0.37$) between different lots of media and reagents.

Table 2. Data for Determination of Ruggedness

Sample	Replicate 1, log(MPN/g)	Replicate 2, log(MPN/g)
1	-0.44	0.17
2	-0.04	0.36
3	0.36	0.36
4	1.96	1.96
5	2.36	1.87
6	1.87	1.96
7	2.36	2.36
8	3.97	3.88
9	4.38	4.66
10	3.88	3.97

3. Precision & Recovery

Precision

Purpose/Method

The difference between the methods results (MPNs) and the reference values should be consistent both between different samples and also when detecting varying concentrations of measurand. The precision of the method tests the consistency of the difference between the reference values and the MPN values found in spiked matrix. This was done by testing ten oyster samples at low, medium, and high concentrations in duplicate to determine the MPN. The MPN and reference data sets were converted into log values. Each MPN was compared to its associate reference value. A nested ANOVA was then performed, with variance components being the sample, and concentrations within the samples (low, medium, and high), and then the error. The ANOVA component of interest was that comparing the concentrations within the samples to the determinations (or error).

Results

The difference between reference values and MPNs can be found below in Table 3. The data shows that there are no significant differences between the concentrations in samples and the determinations within concentrations via a nested ANOVA ($p=0.69$). Additionally, the variance of the method does not exceed the known variance of a 3-tube MPN ($p=0.16$).

Table 3: Data for determining the Precision

Sample	Concentration	Difference	log(reference MPNs)	log(MPNs)
1	low	0.27	0.23	-0.04
1	low	-0.40	0.23	0.63
1	med	-0.55	3.41	3.97
1	med	0.45	3.41	2.96
1	high	-0.24	4.41	4.66
1	high	-0.24	4.41	4.66
2	low	-0.19	0.18	0.36
2	low	0.21	0.18	-0.04
2	med	0.06	3.23	3.17
2	med	-0.13	3.23	3.36
2	high	0.26	4.23	3.97
2	high	0.06	4.23	4.17
3	low	0.27	0.13	-0.14
3	low	0.17	0.13	-0.04
3	med	0.21	3.18	2.96
3	med	-0.28	3.18	3.46
3	high	-0.20	4.18	4.38
3	high	0.21	4.18	3.97
4	low	0.23	0.29	0.06
4	low	0.73	0.29	-0.44
4	med	-0.23	3.13	3.36
4	med	0.17	3.13	2.96

4	high	-0.20	4.13	4.33
4	high	-0.53	4.13	4.66
5	low	-0.13	0.18	0.31
5	low	-0.45	0.18	0.63
5	med	-0.07	3.29	3.36
5	med	-0.07	3.29	3.36
5	high	-0.09	4.29	4.38
5	high	0.32	4.29	3.97
6	low	0.13	0.49	0.36
6	low	0.53	0.49	-0.04
6	med	-0.14	3.49	3.63
6	med	0.32	3.49	3.17
6	high	0.00	4.18	4.17
6	high	0.00	4.18	4.17
7	low	-0.05	0.31	0.36
7	low	0.35	0.31	-0.04
7	med	-0.05	3.31	3.36
7	med	0.35	3.31	2.96
7	high	0.34	4.31	3.97
7	high	0.34	4.31	3.97
8	low	-0.28	0.08	0.36
8	low	0.22	0.08	-0.14
8	med	0.12	3.08	2.96
8	med	-0.28	3.08	3.36
8	high	-0.25	4.08	4.33

8	high	-0.09	4.08	4.17
9	low	0.27	0.44	0.17
9	low	0.12	0.44	0.32
9	med	0.08	3.44	3.36
9	med	0.27	3.44	3.17
9	high	0.06	4.44	4.38
9	high	0.27	4.44	4.17
10	low	0.13	0.30	0.17
10	low	0.34	0.30	-0.04
10	med	-0.06	3.30	3.36
10	med	0.67	3.30	2.63
10	high	0.13	4.30	4.17
10	high	0.33	4.30	3.97

Recovery

Purpose/Method

The amount of analyte recovered should be consistent both between different samples and also when detecting varying concentrations of measurand. The recovery of the method tests the consistency of the analyte recovered via MPNs as compared to the reference values. This was done by testing ten oyster samples at low, medium, and high concentrations in duplicate to determine the MPN. Each MPN and reference value was converted to logs. Each duplicated MPN was averaged and then compared to its associate reference value, in logs. A single-factor ANOVA was then used to compare the recovery at the three concentrations.

Results

The difference between reference values and MPNs can be found below in Table 4. The recovery across all samples and concentrations was found to be 97.44%. There was not found to be significant differences in the recovery at the various concentrations ($p=0.49$).

Table 4: Data for the determination of Recovery

Sample	Concentration	Avg log(Reference) per Conc.	Avg log(MPN) per Conc.
1	low	0.23	0.30
	med	3.41	3.47
	high	4.41	4.66
2	low	0.18	0.16
	med	3.23	3.27
	high	4.23	4.07
3	low	0.13	-0.09
	med	3.18	3.21
	high	4.18	4.17
4	low	0.29	-0.19
	med	3.13	3.16
	high	4.13	4.49
5	low	0.18	0.47
	med	3.29	3.36
	high	4.29	4.17
6	low	0.49	0.16
	med	3.49	3.40
	high	4.18	4.17
7	low	0.31	0.16
	med	3.31	3.16
	high	4.31	3.97
8	low	0.08	0.11
	med	3.08	3.16
	high	4.08	4.25
9	low	0.44	0.24
	med	3.44	3.27
	high	4.44	4.28
10	low	0.30	0.07
	med	3.30	3.00
	high	4.30	4.07

4. Specificity

Purpose/Method

The method should only detect the analyte of interest, even in the presence of interfering

organisms. Specificity refers to the ability of the method to measure only the target organism. One matrix sample was divided into two aliquots. One aliquot was spiked with a low but determinable level of *Vibrio vulnificus* (*Vv*). The other aliquot was spiked with the same level of *Vv* as the first, but also spiked with a high level of potential interfering *Vibrio parahaemolyticus* (*Vp*). Five replicates were performed. Each of the replicates was analyzed by taking the average log MPN and calculating the Specificity Index (SI). A paired *t*-test was used to determine if the average specificity index obtained from the five replicates differed from 1 (significance level = 0.05).

Results

Using the data from Table 5, the average specificity index was 0.99 when in the presence of *Vp*. These values are not significantly different than 1 ($p=0.74$).

Table 5. Data for Determination of Specificity

Replicate	<i>Vv</i> only, log(MPN/g)	<i>Vp</i> + <i>Vv</i> , log(MPN/g)
1	1.63	1.96
2	2.32	2.17
3	1.96	1.63
4	1.96	2.36
5	2.17	2.17

5. Linear Range, Limit of Detection & Limit of Quantification/Sensitivity

Linear Range

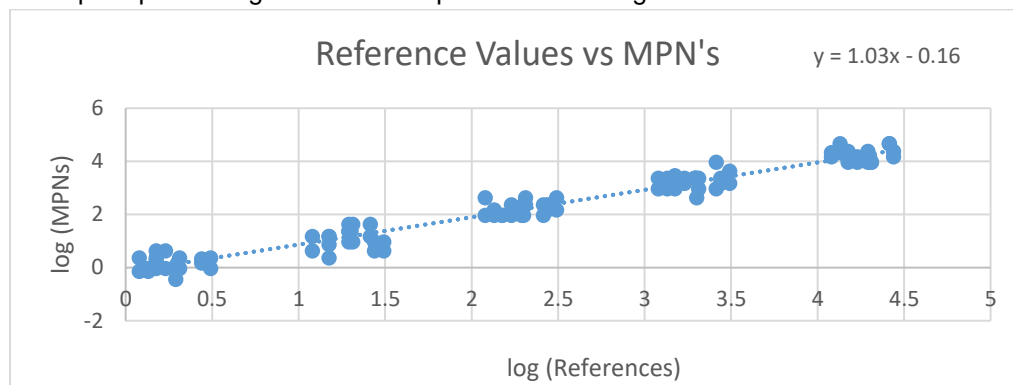
Purpose/Method

The MPN value found should directly correlate to the concentration of analyte within the sample, within the working range of the method. Thus, as the concentration increases, the MPN value should also increase in a linear fashion. Ten Oyster samples were tested at 5 concentration levels, in duplicate. Each MPN was compared to its associate reference value. The relationship between the log(MPN) and log(reference) was then found by obtaining the correlation coefficient by performing a linear regression with log(reference) as the independent variable and log(MPN) as the dependent variable.

Results

The relationship between the MPNs and reference values can be seen in Figure 1 below. The relationship between MPNs and reference values was found to be linear, with a Pearson's *r* of 0.98. The working range used was of concentrations ranging from 10^0 to 10^4 cells/gram.

Figure 1: Graph representing the relationship between the log of reference values and MPNS



Limit of Detection

Purpose/Method

The method should be capable of detecting as little as 1 cell/gram of sample, or 0 cells/gram, in log form. Therefore, it must be determined whether the method can detect one cell per gram of sample. The log(MPN) was compared to the log(reference) of ten oyster samples, spiked at five varying concentrations, in duplicate. This was done by performing a regression analysis on the data and calculating the Limit of Detection by taking the antilog of the intercept. The independent variable was set as log(reference) and the dependent variable was set as log(MPN).

Results

The Limit of Detection was found to be 0.68 cells. The overall regression 95.0% confidence interval was found to be 0.58, which encompasses the Limit of Detection. The 99.99% confidence interval of the intercept was found to be 0.23. These both contain the intercept value of -0.16 within the interval.

Table 7: Data for determination of the Limit of Detection

Sample	Concentration	Log(reference)	Log(MPN)
Sample 1, Rep 1	10 ⁰	0.23	-0.04
	10 ¹	1.41	1.17
	10 ²	2.41	1.96

	10 ³	3.41	3.97
	10 ⁴	4.41	4.66
Sample 1, Rep 2	10 ⁰	0.23	0.63
	10 ¹	1.41	1.63
	10 ²	2.41	2.36
	10 ³	3.41	2.96
	10 ⁴	4.41	4.66
Sample 2, Rep 1	10 ⁰	0.18	0.36
	10 ¹	1.18	1.17
	10 ²	2.23	1.96
	10 ³	3.23	3.17
	10 ⁴	4.23	3.97
Sample 2, Rep 2	10 ⁰	0.18	-0.04
	10 ¹	1.18	0.87
	10 ²	2.23	2.36
	10 ³	3.23	3.36
	10 ⁴	4.23	4.17

Sample 3, Rep 1	10⁰	0.13	-0.14
	10¹	1.29	0.96
	10²	2.29	1.96
	10³	3.18	2.96
	10⁴	4.18	4.38
Sample 3, Rep 2	10⁰	0.13	-0.04
	10¹	1.29	1.36
	10²	2.29	2.17
	10³	3.18	3.46
	10⁴	4.18	3.97
Sample 4, Rep 1	10⁰	0.29	0.06
	10¹	1.29	1.36
	10²	2.13	2.17
	10³	3.13	3.36
	10⁴	4.13	4.33
Sample 4, Rep 2	10⁰	0.29	-0.44
	10¹	1.29	1.63

	10 ²	2.13	1.96
	10 ³	3.13	2.96
	10 ⁴	4.13	4.66
Sample 5, Rep 1	10 ⁰	0.18	0.31
	10 ¹	1.18	0.36
	10 ²	2.18	1.96
	10 ³	3.29	3.36
	10 ⁴	4.29	4.38
Sample 5, Rep 2	10 ⁰	0.18	0.63
	10 ¹	1.18	1.17
	10 ²	2.18	1.96
	10 ³	3.29	3.36
	10 ⁴	4.29	3.97
Sample 6, Rep 1	10 ⁰	0.49	0.36
	10 ⁰	1.49	0.63
	10 ²	2.49	2.63
	10 ³	3.49	3.63

	10 ⁴	4.18	4.17
Sample 6, Rep 2	10 ⁰	0.49	-0.04
	10 ¹	1.49	0.96
	10 ²	2.49	2.17
	10 ³	3.49	3.17
	10 ⁴	4.18	4.17
	Sample 7, Rep 1	10 ⁰	0.31
10 ¹		1.31	0.96
10 ²		2.31	2.36
10 ³		3.31	3.36
10 ⁴		4.31	3.97
Sample 7, Rep 2	10 ⁰	0.31	-0.04
	10 ¹	1.31	1.63
	10 ²	2.31	2.63
	10 ³	3.31	2.96
	10 ⁴	4.31	3.97
Sample 8, Rep 1	10 ⁰	0.08	0.36

	10 ¹	1.08	1.17
	10 ²	2.08	2.63
	10 ³	3.08	2.96
	10 ⁴	4.08	4.33
Sample 8, Rep 2	10 ⁰	0.08	-0.14
	10 ¹	1.08	0.63
	10 ²	2.08	1.96
	10 ³	3.08	3.36
	10 ⁴	4.08	4.17
Sample 9, Rep 1	10 ⁰	0.44	0.17
	10 ¹	1.44	0.96
	10 ²	2.44	2.17
	10 ³	3.44	3.36
	10 ⁴	4.44	4.38
Sample 9, Rep 2	10 ⁰	0.44	0.32
	10 ¹	1.44	0.63
	10 ²	2.44	2.36

	10³	3.44	3.17
	10⁴	4.44	4.17
Sample 10, Rep 1	10⁰	0.30	0.17
	10¹	1.30	1.36
	10²	2.30	1.96
	10³	3.30	3.36
	10⁴	4.30	4.17
Sample 10, Rep 2	10⁰	0.30	-0.04
	10¹	1.30	1.36
	10²	2.30	2.36
	10³	3.30	2.63
	10⁴	4.30	3.97

Limit of Quantification/Sensitivity

Purpose/Method

The quantifiable limit of the method is bounded by the values defined by a 3-tube MPN. In the case that the Limit of Detection is not significantly different than 1 cell/gram, than the Limit of Quantification can be extrapolated using the FDA BAM MPN Calculator.

Results

As the method starts with a low dilution of 1 gram of sample per tube, use of a 3-tube MPN and corresponding dilution ratios will result in the Limit of Quantification/Sensitivity for the method being 0.36 MPN/gram.

Inclusivity

Purpose

To assess the ability of the method to detect a wide range of target strains in various oyster tissues.

Method

Vibrio vulnificus (Vv) strains were grown in APW for 18-24 hours. DNA was extracted using the MagNAPure 96 instrument and PCR performed using the QuantStudio Dx Real-Time PCR System. The strains that make up the inclusivity panel were obtained from the Center for Disease Control and Prevention (CDC). The primers and probes utilized in this method have been tested against DNA extracts from the isolates listed in Table 1 below. The *vvhA* marker was either confirmed by the CDC or ATCC paperwork.

Results

Primer / Probe Sensitivity

Sensitivity= (# of true positives/ (# of true positives + # of false negatives))

vvhA sensitivity = 21/21 = **100% *vvhA* sensitivity**

The primers and probes utilized in this method demonstrates 100% inclusivity. See Table 1 (Inclusivity Panel) for the breakdown of strains, source, targets tested, and its corresponding result.

Table 1. Inclusivity panel with results by target. Targets not tested or confirmed are blocked out.

Strain	Source	tlh	tdh	trh	orf8	vvhA
06-2410	06-2410 (CDC)					+
06-2450	06-2450 (CDC)					+
07-2405	07-2405 (CDC)					+
08-2468	08-2468 (CDC)					+
08-2470	08-2470 (CDC)					+
08-2472	08-2472 (CDC)					+
08-2485	08-2485 (CDC)					+
1831-81	1831-81 (CDC)					+
2009V-1002	2009V-1002 (CDC)					+
2009V-1055	2009V-1055 (CDC)					+
2010V-1021	2010V-1021 (CDC)					+
209V-1035	209V-1035 (CDC)					+
2431-04	2431-04 (CDC)					+
2473-85	2473-85 (CDC)					+
2492-88	2492-88 (CDC)					+

2809-78	2809-78 (CDC)					+
430-79	430-79 (CDC)					+
AM38622	AM38622 (CDC)					+
AM38623	AM38623 (CDC)					+
27562	27562					+
29307	29307					+
Total Confirmed Isolated		0	0	0	0	21

Exclusivity

Purpose

To demonstrate the ability of the method to distinguish the targeted analyte from other potentially cross-reactive non-target strains that could possibly contaminate shellfish.

Method

All organisms were inoculated into APW and incubated for 18-24 hours. DNA was extracted using the MagNAPure 96 instrument and PCR performed using the QuantStudio Dx Real-Time PCR System. All strains were obtained from the Center for Disease Control and Prevention (CDC) and American Type Culture Collection (ATCC).

Results

Primer / Probe Specificity

Specificity = (# of true negative/ (# of true negatives + # of false positives))

vvhA specificity = 28/28 = **100% *vvhA* Specificity**

The *vvhA* primers and probes utilized in this method demonstrate 100% exclusivity. None of the exclusivity panel had detection of *vvhA*. See Table 2 (Exclusivity Panel) for the breakdown of strains, source, targets tested, and its corresponding result.

Table 2. Exclusivity panel with results by target. Targets not tested or confirmed are blocked out.

Organism	ATCC #	<i>tlh</i>	<i>tdh</i>	<i>trh</i>	<i>orf8</i>	<i>vvhA</i>
<i>E. coli</i>	25922	-	-	-	-	-
<i>G. hollisae</i>	33564	-	-	-	-	-
<i>K. pneumoniae</i>	33495	-	-	-	-	-
<i>P. aeruginosa</i>	33495	-	-	-	-	-
<i>S. aureus</i>	10145	-	-	-	-	-
<i>S. sonnei</i>	25925	-	-	-	-	-

<i>S. typhimurium</i>	9290	-	-	-	-	-
<i>V. aestuarians</i>	35048	-	-	-	-	-
<i>V. alginolyticus</i>	17749	-	-	-	-	-
<i>V. alginosus</i>	14390	-	-	-	-	-
<i>V. campbellii</i>	25920	-	-	-	-	-
<i>V. cholerae</i>	39050	-	-	-	-	-
<i>V. cincinnatiensis</i>	35912	-	-	-	-	-
<i>V. furnissii</i>	33813	-	-	-	-	-
<i>V. marinagilis</i>	14398	-	-	-	-	-
<i>V. marinofulvus</i>	14395	-	-	-	-	-
<i>V. marinovulgaris</i>	14394	-	-	-	-	-
<i>V. metschnikovii</i>	700040	-	-	-	-	-
<i>V. mimicus</i>	33653	-	-	-	-	-
<i>V. natriegens</i>	14048	-	-	-	-	-
<i>V. nereis</i>	25917	-	-	-	-	-
<i>V. nigripulchritudo</i>	27043	-	-	-	-	-
<i>V. ponticus</i>	14391	-	-	-	-	-
<i>V. proteolyticus</i>	15338	-	-	-	-	-
<i>V. splendidus</i>	33789	-	-	-	-	-
<i>V. tubiashii</i>	19106	-	-	-	-	-
<i>A. trota</i>	2013V-1197 (CDC)	-	-	-	-	-
<i>A. veronii</i>	N/A (CDC)	-	-	-	-	-
Total Strains		28				

**PUBLIC HEALTH SERVICE
U.S. FOOD AND DRUG ADMINISTRATION
OFFICE OF FOOD SAFETY
SHELLFISH AND AQUACULTURE POLICY BRANCH
5100 PAINT BRANCH PARKWAY
COLLEGE PARK, MD 20740-3835
TEL. 240- 402-2151/2055/4960 FAX 301-436-2601**

SHELLFISH LABORATORY EVALUATION CHECKLIST

LABORATORY:

ADDRESS:

TELEPHONE:

FAX:

EMAIL:

DATE OF EVALUATION:

DATE OF REPORT:

LAST EVALUATION:

LABORATORY REPRESENTED BY:

TITLE:

LABORATORY EVALUATION OFFICER:

SHELLFISH SPECIALIST:

REGION:

OTHER OFFICIALS PRESENT:

TITLE:

Items which do not conform are noted by:

Conformity it noted by a “√”

C- Critical K - Key O - Other NA- Not Applicable

Check the applicable analytical methods:

- | | |
|--------------------------|--|
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Seawater (APHA)[PART II] |
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Seawater using MA-1 [PART II] |
| <input type="checkbox"/> | Membrane Filtration Technique for Seawater using mTEC [PART II] |
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Shellfish Meats (APHA)[PART III] |
| <input type="checkbox"/> | Standard Plate Count for Shellfish Meats [PART III] |
| <input type="checkbox"/> | Elevated Temperature Coliform Plate Method for Shellfish Meats [PART III] |
| <input type="checkbox"/> | Male Specific Coliphage for Soft-shelled Clams and American Oysters [PART III] |

PART 1 - QUALITY ASSURANCE

CODE	REF.	ITEM	
K	8, 11	1.1 Quality Assurance (QA) Plan	
		<input type="checkbox"/>	1.1.1 Written Plan (Check those items which apply.)
		<input type="checkbox"/>	a. Organization of the laboratory.
		<input type="checkbox"/>	b. Staff training requirements.
		<input type="checkbox"/>	c. Standard operating procedures.
		<input type="checkbox"/>	d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		<input type="checkbox"/>	e. Laboratory safety.
		<input type="checkbox"/>	f. Internal performance assessment.
C	8	<input type="checkbox"/>	1.1.2 QA Plan Implemented.
K	11	<input type="checkbox"/>	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s)_____
1.2 Educational/Experience Requirements			
C	State's Human Resources Department	<input type="checkbox"/>	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	<input type="checkbox"/>	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three months of experience in laboratory sciences.
1.3 Work Area			
O	8,11	<input type="checkbox"/>	1.3.1 Adequate for workload and storage.
K	11	<input type="checkbox"/>	1.3.2 Clean, well-lighted.
K	11	<input type="checkbox"/>	1.3.3 Adequate temperature control.
O	11	<input type="checkbox"/>	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	<input type="checkbox"/>	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
1.4 Laboratory Equipment			
O	9	<input type="checkbox"/>	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	<input type="checkbox"/>	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	<input type="checkbox"/>	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	<input type="checkbox"/>	1.4.4 pH meter is calibrated daily or with each use Results are recorded and records maintained.
K	11	<input type="checkbox"/>	1.4.5 A minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	8,15	<input type="checkbox"/>	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt

				procedure or through determination of the slope. <i>(Circle the method used.)</i>
K	9	<input type="checkbox"/>	1.4.7	Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13	<input type="checkbox"/>	1.4.8	Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11	<input type="checkbox"/>	1.4.9	Refrigerator temperature(s) are monitored at least once daily on workdays. Results are recorded and records maintained.
K	1	<input type="checkbox"/>	1.4.10	Refrigerator temperature is maintained at 0 to 4°C.
C	9	<input type="checkbox"/>	1.4.11	The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11	<input type="checkbox"/>	1.4.12	Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9	<input type="checkbox"/>	1.4.13	Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11	<input type="checkbox"/>	1.4.14	Temperature of the waterbath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9	<input type="checkbox"/>	1.4.15	The thermometers used in the waterbath are graduated in at least 0.1°C increments.
C	13	<input type="checkbox"/>	1.4.16	The waterbath has adequate capacity for workload.
K	9	<input type="checkbox"/>	1.4.17	The level of water in the waterbath covers the level of liquid in the incubating tubes.
K	8, 11	<input type="checkbox"/>	1.4.18	Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4	<input type="checkbox"/>	1.4.19	All working thermometers are appropriately immersed.
C	29	<input type="checkbox"/>	1.4.20	Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	11	<input type="checkbox"/>	1.4.21	A mercury-in-glass standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9	<input type="checkbox"/>	1.4.22	Standards thermometers are checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination_____.
C	29	<input type="checkbox"/>	1.4.23	Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of ≤ ±0.05°C are used as the laboratory standards thermometer. <i>(Circle the thermometer type used.)</i>
K	13	<input type="checkbox"/>	1.4.24	Incubator and waterbath working thermometers are checked annually against the standards thermometer at the temperatures at which they are used. Results are recorded and records maintained.
O	11	<input type="checkbox"/>	1.4.25	Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing				
O	9	<input type="checkbox"/>	1.5.1	Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding materials.
K	9	<input type="checkbox"/>	1.5.2	Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.
K	9	<input type="checkbox"/>	1.5.3	Sample containers are made of glass or some other inert material.
O	9	<input type="checkbox"/>	1.5.4	Dilution bottles and tubes are made of borosilicate glass or plastic and closed with rubber stoppers, caps or screw caps with nontoxic liners.

K	9	<input type="checkbox"/>	1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9	<input type="checkbox"/>	1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1mL used to deliver 0.1 mL aliquots.
K	9	<input type="checkbox"/>	1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9	<input type="checkbox"/>	1.5.8	In washing reusable pipettes, a succession of at least three fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2	<input type="checkbox"/>	1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11	<input type="checkbox"/>	1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination				
K	9	<input type="checkbox"/>	1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8	<input type="checkbox"/>	1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	11, 30	<input type="checkbox"/>	1.6.3	The autoclave provides a sterilizing temperature of 121± 2°C as determined for each load using a calibrated maximum registering thermometer. As an alternative, an appropriate temperature monitoring device is used in place of the maximum registering thermometer when these are unavailable due to the ban on mercury.
K	11	<input type="checkbox"/>	1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16	<input type="checkbox"/>	1.6.5	The autoclave standards thermometer is checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory; or, is checked in-house at the steam point (100°C) if it has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination_____
K	1	<input type="checkbox"/>	1.6.6	Working autoclave thermometers are checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11	<input type="checkbox"/>	1.6.7	Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11	<input type="checkbox"/>	1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13	<input type="checkbox"/>	1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11	<input type="checkbox"/>	1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180°C.
K	9	<input type="checkbox"/>	1.6.11	A thermometer capable of determining temperatures accurately in the range of 160 to 180°C is used to monitor the operation of the hot-air sterilizing oven
K	13	<input type="checkbox"/>	1.6.12	Records of temperatures and exposure times are maintained for the operation of the hot-air sterilizing oven during use.
K	11	<input type="checkbox"/>	1.6.13	Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Records are maintained.

K	11	<input type="checkbox"/>	1.6.14	Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C.
C	1	<input type="checkbox"/>	1.6.15	The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1	<input type="checkbox"/>	1.6.16	The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9	<input type="checkbox"/>	1.6.17	Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9	<input type="checkbox"/>	1.6.18	Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2 hours.
C	2	<input type="checkbox"/>	1.6.19	The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2	<input type="checkbox"/>	1.6.20	The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18	<input type="checkbox"/>	1.6.21	Hardwood applicator transfer sticks are properly sterilized. Method of sterilization _____
C	2	<input type="checkbox"/>	1.6.22	The sterility of the hardwood applicator transfer sticks is checked routinely. Results are recorded and the records maintained.
O	13	<input type="checkbox"/>	1.6.23	Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation				
K	3, 5	<input type="checkbox"/>	1.7.1	Media is commercially dehydrated except in the case of medium A-1 which must be prepared from the individual components and modified MacConkey agar which may be prepared from its components.
K	11	<input type="checkbox"/>	1.7.2	Media is prepared according to manufacturer's instructions.
O	11	<input type="checkbox"/>	1.7.3	Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11	<input type="checkbox"/>	1.7.4	Dehydrated media are labeled with date of receipt and date opened.
C	12	<input type="checkbox"/>	1.7.5	Caked or expired media or media components are discarded.
C	133	<input type="checkbox"/>	1.7.6	Reagent water is distilled or deionized (circle appropriate choice), tested monthly and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) <u>for mixed-bed deionizers resistivity (measured in-line) should be >10 megohm-cm at 25°C</u> or is less than 2.0 µSiemens/cm conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11	<input type="checkbox"/>	1.7.7	Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level (< 0.1 mg/L). Results are recorded and the records maintained. Specify method of determination _____.
K	11	<input type="checkbox"/>	1.7.8	Reagent water contains <100 CFU/mL as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11	<input type="checkbox"/>	1.7.9	Media prepared from commercial dehydrated components are sterilized according to the manufacturer's instructions.
K	9	<input type="checkbox"/>	1.7.10	The volume and concentration of media in the tube are suitable for the amount of sample inoculated.
C	11	<input type="checkbox"/>	1.7.11	Total time of exposure of sugar broths to autoclave temperatures does not exceed 45 minutes.
C	1	<input type="checkbox"/>	1.7.12	Media sterility is determined for each load sterilized. Results are recorded and the records maintained.
C	1	<input type="checkbox"/>	1.7.13	Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is

				made from its individual components.
O	9	<input type="checkbox"/>	1.7.14	Sterile phosphate buffered dilution water is used as the sample diluent.
K	11	<input type="checkbox"/>	1.7.15	The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media				
K	9	<input type="checkbox"/>	1.8.1	Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11	<input type="checkbox"/>	1.8.2	Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13	<input type="checkbox"/>	1.8.3	Stored media are labeled with the storage expiration date or the sterilization date.
K	9	<input type="checkbox"/>	1.8.4	Storage of prepared culture media at room temperature does not exceed 7 days.
K	2	<input type="checkbox"/>	1.8.5	Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed 1 month.
K	11	<input type="checkbox"/>	1.8.6	Storage under refrigeration of prepared culture media with screw-cap closures does not exceed 3 months.
K	17	<input type="checkbox"/>	1.8.7	All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES				
2.1 Collection and Transportation of Samples				
C	11	<input type="checkbox"/>	2.1.1	Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1	<input type="checkbox"/>	2.1.2	Samples are identified with collectors name, harvest area, sampling station, time and date of collection.
C	9	<input type="checkbox"/>	2.1.3	Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1	<input type="checkbox"/>	2.1.4	A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9	<input type="checkbox"/>	2.1.5	Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.
2.2 Bacteriological Examination of Seawater by the APHA MPN				
C	9	<input type="checkbox"/>	2.2.1	Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
C	2	<input type="checkbox"/>	2.2.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	9	<input type="checkbox"/>	2.2.3	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9	<input type="checkbox"/>	2.2.4	In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	<input type="checkbox"/>	2.2.5	In a single dilution series not less than 12 tubes are used (for depuration at least 5 tubes are used).
C	6	<input type="checkbox"/>	2.2.6	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____

			Range of MPN _____
			Strength of media used _____
K	9	<input type="checkbox"/>	2.2.7 Inoculated tubes are incubated in air at 35 ± 0.5°C.
C	2	<input type="checkbox"/>	2.2.8 Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained.
			Positive process control _____ Negative process control _____
K	9	<input type="checkbox"/>	2.2.9 Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
			2.3 Confirmed Test for Seawater by APHA MPN
C	9	<input type="checkbox"/>	2.3.1 Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9	<input type="checkbox"/>	2.3.2 EC medium is used as the confirmatory medium for fecal coliforms.
C	2	<input type="checkbox"/>	2.3.3 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
			Positive productivity control _____ Negative productivity control _____
K	9, 11	<input type="checkbox"/>	2.3.4 Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. <i>(Circle the method of transfer.)</i>
C	9	<input type="checkbox"/>	2.3.5 BGB tubes are incubated at 35 ± 0.5°C.
K	9	<input type="checkbox"/>	2.3.6 BGB tubes are read after 48 ± 3 hours of incubation.
C	9	<input type="checkbox"/>	2.3.7 EC tubes are incubated in a circulating waterbath maintained at 44.5 ± 0.2°C.
C	9	<input type="checkbox"/>	2.3.8 EC tubes are read after 24 ± 2 hours of incubation.
C	9	<input type="checkbox"/>	2.3.9 The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
			2.4 Computation of Results – APHA MPN
K	9	<input type="checkbox"/>	2.4.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7	<input type="checkbox"/>	2.4.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	<input type="checkbox"/>	2.4.3 Results are reported as MPN/100 mL of sample.
			2.5 Bacteriological Examination of Seawater by the MA-1 Method
C	5	<input type="checkbox"/>	2.5.1 A-1 medium complete is used in the analysis.
C	2, 31	<input type="checkbox"/>	2.5.2 A-1 medium without salicin is used in the analysis. Comparability testing supports use of A-1 medium without salicin. Study records are available.
C	5	<input type="checkbox"/>	2.5.3 A-1 medium sterilized for 10 minutes at 121°C.
C	2	<input type="checkbox"/>	2.5.4 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
			Positive productivity control _____ Negative productivity control _____
C	9	<input type="checkbox"/>	2.5.5 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9	<input type="checkbox"/>	2.5.6 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	<input type="checkbox"/>	2.5.7 In a single dilution series at least 12 tubes are used.

C	6	<input type="checkbox"/>	2.5.8	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	<input type="checkbox"/>	2.5.9	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and waterbath incubation Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	<input type="checkbox"/>	2.5.10	Inoculated tubes are placed in an air incubator at $35 \pm 0.5^\circ\text{C}$ for 3 ± 0.5 hours of resuscitation.
C	5	<input type="checkbox"/>	2.5.11	After 3 ± 0.5 hours resuscitation at 35°C , inoculated tubes are incubated at $44.5 \pm 0.2^\circ\text{C}$ in a circulating waterbath for the remainder of the 24 ± 2 hours.
C	5	<input type="checkbox"/>	2.5.12	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.6 Computation of Results – APHA MPN				
K	9	<input type="checkbox"/>	2.6.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	<input type="checkbox"/>	2.6.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	<input type="checkbox"/>	2.6.3	Results are reported as MPN/100 mL of sample.
2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment				
C	23, 24	<input type="checkbox"/>	2.7.1	When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at $44.5 \pm 0.5^\circ\text{C}$ under any loading capacity.
C	23	<input type="checkbox"/>	2.7.2	When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	<input type="checkbox"/>	2.7.3	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	<input type="checkbox"/>	2.7.4	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	<input type="checkbox"/>	2.7.5	Colonies are counted with the aid of magnification.
C	11, 23	<input type="checkbox"/>	2.7.6	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of $0.45 \mu\text{m}$ and certified by the manufacturer for fecal coliform analyses.
C	2	<input type="checkbox"/>	2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
C	2	<input type="checkbox"/>	2.7.8	When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison testing implemented. The results are recorded and this record is maintained.
K	2, 11	<input type="checkbox"/>	2.7.9	New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2	<input type="checkbox"/>	2.7.10	The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2	<input type="checkbox"/>	2.7.11	Membrane filters which are beyond their expiration date are not used.
O	11	<input type="checkbox"/>	2.7.12	Forceps tips are clean.
O	11	<input type="checkbox"/>	2.7.13	Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.

K	11	<input type="checkbox"/>	2.7.14	Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11	<input type="checkbox"/>	2.7.15	If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11	<input type="checkbox"/>	2.7.16	Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11	<input type="checkbox"/>	2.7.17	Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26	<input type="checkbox"/>	2.7.18	A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11	<input type="checkbox"/>	2.7.19	The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2	<input type="checkbox"/>	2.7.20	Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
2.8 Media Preparation and Storage – MF using mTEC Agar				
K	11	<input type="checkbox"/>	2.8.1	Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
C	11	<input type="checkbox"/>	2.8.2	The phosphate buffered saline is properly sterilized.
K	23	<input type="checkbox"/>	2.8.3	A sufficient amount of medium (4-5 mL) is used in each plate.
O	11	<input type="checkbox"/>	2.8.4	Refrigerated prepared plates are stored for no more than 2 weeks in sealed plastic bags or containers to minimize evaporation.
2.9 Sample Analyses - MF using mTEC Agar				
C	24	<input type="checkbox"/>	2.9.1	mTEC agar is used.
C	2	<input type="checkbox"/>	2.9.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	23	<input type="checkbox"/>	2.9.3	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration.
C	23	<input type="checkbox"/>	2.9.4	The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25	<input type="checkbox"/>	2.9.5	Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23	<input type="checkbox"/>	2.9.6	Sample volumes are filtered under vacuum.
K	26	<input type="checkbox"/>	2.9.7	The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26	<input type="checkbox"/>	2.9.8	The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.
C	23	<input type="checkbox"/>	2.9.9	The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11	<input type="checkbox"/>	2.9.10	Blanks are run at the beginning of filtration, after every 10th aliquot and at the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).
C	2, 11	<input type="checkbox"/>	2.9.11	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24	<input type="checkbox"/>	2.9.12	Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at 35 + 0.5°C for 2 hours of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5°C for 24 ± 2 hours.

C	11, 23, 24	<input type="checkbox"/>	2.9.13	After 2 hours of resuscitation at 35°C, the watertight, tightly sealed containers are transferred to a circulating waterbath at 44.5 + 0.2°C, submerged completely and incubated for 22-24 hours.
2.10 Computation of Results - MF using mTEC Agar				
C	23	<input type="checkbox"/>	2.10.1	All yellow, yellow-green or yellow-brown colonies are counted.
C	23	<input type="checkbox"/>	2.10.2	Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as >80 x 100/the volume of sample filtered.
C	2, 11, 23	<input type="checkbox"/>	2.10.3	When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11	<input type="checkbox"/>	2.10.4	The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11	<input type="checkbox"/>	2.10.5	Results are reported as CFU/100 mL of sample.
PART III - SHELLFISH SAMPLES				
3.1 Collection and Transportation of Samples				
C	9	<input type="checkbox"/>	3.1.1	A representative sample of shellstock is collected.
K	9	<input type="checkbox"/>	3.1.2	Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9	<input type="checkbox"/>	3.1.3	Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9	<input type="checkbox"/>	3.1.4	Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1	<input type="checkbox"/>	3.1.5	Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
3.2 Preparation of Shellfish for Examination				
K	2,11	<input type="checkbox"/>	3.2.1	Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2	<input type="checkbox"/>	3.2.2	Blades of shucking knives are not corroded.
O	9	<input type="checkbox"/>	3.2.3	The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	<input type="checkbox"/>	3.2.4	The faucet used for rinsing the shellstock does not contain an aerator.
K	9	<input type="checkbox"/>	3.2.5	Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	<input type="checkbox"/>	3.2.6	Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	9	<input type="checkbox"/>	3.2.7	Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9	<input type="checkbox"/>	3.2.8	Shellstock are not shucked directly through the hinge.
C	9	<input type="checkbox"/>	3.2.9	Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	<input type="checkbox"/>	3.2.10	At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.
K	9	<input type="checkbox"/>	3.2.11	A representative sample of at least 12 shellfish is used for the analysis.
K	2	<input type="checkbox"/>	3.2.12	The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
O	9	<input type="checkbox"/>	3.2.13	Sterile phosphate buffered dilution water is used as the sample diluent.

C	9	<input type="checkbox"/>	3.2.14	Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9	<input type="checkbox"/>	3.2.15	APHA <i>Recommended Procedures for the Examination of Sea Water And Shellfish</i> , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA				
C	9	<input type="checkbox"/>	3.3.1	Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. <i>(Circle the medium used.)</i>
C	2	<input type="checkbox"/>	3.3.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9	<input type="checkbox"/>	3.3.3	Immediately (within 2 minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9	<input type="checkbox"/>	3.3.4	No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
C	9	<input type="checkbox"/>	3.3.5	Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6	<input type="checkbox"/>	3.3.6	In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	<input type="checkbox"/>	3.3.7	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9	<input type="checkbox"/>	3.3.8	Inoculated media are incubated at $35 \pm 0.5^{\circ}\text{C}$.
K	10	<input type="checkbox"/>	3.3.9	Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA				
C	9	<input type="checkbox"/>	3.4.1	EC medium is used as the confirmatory medium.
C	2	<input type="checkbox"/>	3.4.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11	<input type="checkbox"/>	3.4.3	Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. <i>(Circle the method of transfer.)</i>
C	9	<input type="checkbox"/>	3.4.4	EC tubes are incubated in a circulating waterbath at $44.5 \pm 0.2^{\circ}\text{C}$
K	9	<input type="checkbox"/>	3.4.5	EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9	<input type="checkbox"/>	3.4.6	The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.
3.5 Computation of Results for MPN Analyses				
K	9	<input type="checkbox"/>	3.5.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.
K	7	<input type="checkbox"/>	3.5.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9	<input type="checkbox"/>	3.5.3	Results are reported as MPN/100 grams of sample.

3.6 Standard Plate Count Method			
O	20	<input type="checkbox"/>	3.6.1 A standard plate count (SPC) analysis may be performed in conjunction with the analysis for fecal coliform organisms.
K	9	<input type="checkbox"/>	3.6.2 In the standard plate count procedure at least four plates are used, duplicates of two dilutions. One of the dilutions should produce colonies of 30 to 300 per plate.
K	2	<input type="checkbox"/>	3.6.3 Fifteen to 20 mL of tempered sterile plate count agar is used per plate.
C	9	<input type="checkbox"/>	3.6.4 Agar tempering bath maintains the agar at 44-46°C.
C	9	<input type="checkbox"/>	3.6.5 An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9	<input type="checkbox"/>	3.6.6 Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in 7 seconds) before plating.
C	9	<input type="checkbox"/>	3.6.7 Not more than 1 mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11	<input type="checkbox"/>	3.6.8 Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9,21	<input type="checkbox"/>	3.6.9 Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four high.
K	9	<input type="checkbox"/>	3.6.10 Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1	<input type="checkbox"/>	3.6.11 A hand tally or its equivalent is used for accuracy in counting.
3.7 Computation of Results -SPC			
K	9	<input type="checkbox"/>	3.7.1 Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19	<input type="checkbox"/>	3.7.2 Colony counts are reported as CFU/g of sample.
3.8 Bacteriological Analysis of Shellfish Using the ETCP			
C	2,3	<input type="checkbox"/>	3.8.1 Prepared modified MacConkey agar is used on the day that it is made.
K	3	<input type="checkbox"/>	3.8.2 Double strength modified MacConkey agar is used.
C	3	<input type="checkbox"/>	3.8.3 Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3	<input type="checkbox"/>	3.8.4 Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50°C until used.
K	2, 3	<input type="checkbox"/>	3.8.5 Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3	<input type="checkbox"/>	3.8.6 The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9	<input type="checkbox"/>	3.8.7 The sample homogenate is cultured within 2 minutes of blending.
C	2,3	<input type="checkbox"/>	3.8.8 Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.
K	3	<input type="checkbox"/>	3.8.9 Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2,3, 22	<input type="checkbox"/>	3.8.10 The container is gently swirled or slowly inverted once to mix the contents, which are subsequently distributed uniformly over six plates.
C	1	<input type="checkbox"/>	3.8.11 Media and diluent sterility are determined with each use. Results are recorded and the records maintained.
C	1	<input type="checkbox"/>	3.8.12 Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture _____ Negative control culture _____
C	3, 13	<input type="checkbox"/>	3.8.13 When solidified, the plates are placed inverted into an air incubator at 45.5 ± 0.5°C for 18 to 30 hours of incubation.
C	2	<input type="checkbox"/>	3.8.14 Plates are stacked no more than three high in the incubator.

C	2	<input type="checkbox"/>	3.8.15	Appropriately diluted pour plated process control cultures <i>accompany each set of samples throughout incubation</i> . The results are recorded and the records maintained. Positive process control _____ Negative process control _____
3.9 Computation of Results - ETCP				
K	11	<input type="checkbox"/>	3.9.1	Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1	<input type="checkbox"/>	3.9.2	A hand tally or its equivalent is used to aid in counting.
C	3, 6	<input type="checkbox"/>	3.9.3	All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3	<input type="checkbox"/>	3.9.4	Results are reported as CFU/100 grams of sample.
Bacteriological Examination of Soft-shelled Clams and American Oysters for Male Specific Coliphage (MSC)				
3.10 MSC Equipment and Supplies				
K	30	<input type="checkbox"/>	3.10.1	Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold 100 – 125 mL.
C	27, 28	<input type="checkbox"/>	3.10.2	The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	9	<input type="checkbox"/>	3.10.3	The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
C	27, 28	<input type="checkbox"/>	3.10.4	Sterile 0.22 µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1	<input type="checkbox"/>	3.10.5	The sterility of each lot of pre-sterilized syringes and syringe filters is determined. Results are recorded and records maintained.
K	1	<input type="checkbox"/>	3.10.6	The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28	<input type="checkbox"/>	3.10.7	The balance used provides a sensitivity of at least mg (0.01g).
C	27, 28	<input type="checkbox"/>	3.10.8	The temperature of the incubator used is maintained at 36 ± 1°C.
C	28	<input type="checkbox"/>	3.10.9	Sterile disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
3.11 MSC Media Preparation				
K	28	<input type="checkbox"/>	3.11.1	Media preparation and sterilization is according to the validated method.
K	27, 28	<input type="checkbox"/>	3.11.2	Bottom agar, double strength soft agar and growth broth are prepared from their individual components.
K	27, 28	<input type="checkbox"/>	3.11.3	Soft agar is prepared double strength in volumes of 2.5 mL.
C	27, 28	<input type="checkbox"/>	3.11.4	The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
O	27, 28	<input type="checkbox"/>	3.11.5	Storage of the bottom agar under refrigeration does not exceed 1 month.
K	27, 28	<input type="checkbox"/>	3.11.6	Unsterilized soft agar is stored at -20 °C -15C for up to 3 months.
K	27, 28	<input type="checkbox"/>	3.11.7	The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.
K	27, 28	<input type="checkbox"/>	3.11.8	Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	27, 28	<input type="checkbox"/>	3.11.9	Bottom agar plates are allowed to reach room temperature before use.
3.12 Preparation of the Soft-Shelled Clams and American Oysters for MSC Analysis				
K	2,11	<input type="checkbox"/>	3.12.1	Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2	<input type="checkbox"/>	3.12.2	The blades of shucking knives are not corroded.
O	9	<input type="checkbox"/>	3.12.3	The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	<input type="checkbox"/>	3.12.4	The faucet used for rinsing the shellfish does not contain an aerator.

K	9	<input type="checkbox"/>	3.12.5	The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	<input type="checkbox"/>	3.12.6	The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9	<input type="checkbox"/>	3.12.7	Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9	<input type="checkbox"/>	3.12.8	Shellfish are not shucked through the hinge.
C	9	<input type="checkbox"/>	3.12.9	The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	<input type="checkbox"/>	3.12.10	A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19	<input type="checkbox"/>	3.12.11	The sample is weighed to the nearest 0.1 gram.
3.13 MSC Sample Analysis				
C	28	<input type="checkbox"/>	3.13.1	<i>E.coli Famp ATCC 700891</i> is the bacterial host strain used in this procedure.
K	27, 28	<input type="checkbox"/>	3.13.2	Host cell growth broth is tempered at $36 \pm 1^\circ\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	27, 28	<input type="checkbox"/>	3.13.3	Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^\circ\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	27, 28	<input type="checkbox"/>	3.13.4	After inoculation, the host cell growth broth culture is not shaken.
C	28	<input type="checkbox"/>	3.13.5	A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.
C	28	<input type="checkbox"/>	3.13.6	The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28	<input type="checkbox"/>	3.13.7	The elution mixture is homogenized at high speed for 180 seconds.
C	28	<input type="checkbox"/>	3.13.8	Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
C	28	<input type="checkbox"/>	3.13.9	The homogenized elution mixture is centrifuged for 15 minutes at 9000 x g at 4°C.
C	27, 28	<input type="checkbox"/>	3.13.10	The supernatant is pipetted off, weighed and the weight recorded.
C	27, 28	<input type="checkbox"/>	3.13.11	The supernatant is allowed to warm to room temperature about 20 to 30 minutes.
K	27, 28	<input type="checkbox"/>	3.13.12	The autoclaved soft agar is tempered and held at $51 \pm 1^\circ\text{C}$ throughout the period of sample analysis.
K	27, 28	<input type="checkbox"/>	3.13.13	Two hundred microliters (0.2 mL) of log phase host strain <i>E coli</i> is added to the tempering soft agar immediately prior to adding the sample supernatant.
K	27, 28	<input type="checkbox"/>	3.13.14	The sample supernatant is shaken or vortexed before being added to the tempering soft agar.
C	27, 28	<input type="checkbox"/>	3.13.15	2.5 mL of sample supernatant is added to each tube of tempering soft agar.
C	27, 28	<input type="checkbox"/>	3.13.16	The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28	<input type="checkbox"/>	3.13.17	The soft agar/sample supernatant/host cell mixture is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.
C	28	<input type="checkbox"/>	3.13.18	Ten (10) plates are used, 2.5 mL per plate for a total of 25 mL of supernatant analyzed per sample.
K	27, 28	<input type="checkbox"/>	3.13.19	Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are recorded and records maintained. Positive control _____
K	27, 28	<input type="checkbox"/>	3.13.20	Growth broth is used as the negative control or blank.
K	27, 28	<input type="checkbox"/>	3.13.21	Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2	<input type="checkbox"/>	3.13.22	A negative control plate is plated at the beginning and end of each set of samples analyzed.

K	27, 28	<input type="checkbox"/>	3.13.23 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28	<input type="checkbox"/>	3.13.24 All plates are incubated at 36 ± 1°C for 18 ± 2 hours.
3.14 Computation of Results - MSC			
C	27	<input type="checkbox"/>	3.14.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 32	<input type="checkbox"/>	3.14.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten plates, the count is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/ 100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as > 20,000 PFU/100 grams.
K	28	<input type="checkbox"/>	3.14.3 The formula used for determining the density of MSC in PFU/100 grams is: (0.364) (N) (Ws), where N = total number of plaques counted on all 10 plates and Ws = weight of the supernatant used.
O	9	<input type="checkbox"/>	3.14.4 The MSC count is rounded off conventionally to give a whole number.

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LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
MICROBIOLOGICAL COMPONENT: (Part I-III)	
A. Results	
Total # of Critical (C) Nonconformities in Parts I-III	_____
Total # of Key (K) Nonconformities in Parts I-III	_____
Total # of Critical, Key and Other (O)	_____
Nonconformities in Parts I-III	
B. Criteria for Determining Laboratory Status of the Microbiological Component:	
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <p style="margin-left: 40px;">a. The total # of Critical nonconformities is ≥ 4 or</p> <p style="margin-left: 40px;">b. The total # of Key nonconformities is ≥ 13 or</p> <p style="margin-left: 40px;">c. The total # of Critical, Key and Other is ≥ 18</p> <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but ≤ 3.</p>	
C. Laboratory Status (<i>circle appropriate</i>)	
Does Not Conform	Provisionally Conforms
Conforms	
Acknowledgment by Laboratory Director/Supervisor:	
All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____.	
Laboratory Signature: _____	Date: _____
LEO Signature: _____	Date: _____

NSSP Form LAB-100 Microbiology Rev. October 2015

**PUBLIC HEALTH SERVICE
U.S. FOOD AND DRUG ADMINISTRATION
OFFICE OF FOOD SAFETY
SHELLFISH AND AQUACULTURE POLICY BRANCH
5100 PAINT BRANCH PARKWAY
COLLEGE PARK, MD 20740-3835
TEL. 240- 402-2151/2055/4960 FAX 301-436-2601**

SHELLFISH LABORATORY EVALUATION CHECKLIST

LABORATORY:

ADDRESS:

TELEPHONE:

FAX:

EMAIL:

DATE OF EVALUATION:

DATE OF REPORT:

LAST EVALUATION:

LABORATORY REPRESENTED BY:

TITLE:

LABORATORY EVALUATION OFFICER:

SHELLFISH SPECIALIST:

REGION:

OTHER OFFICIALS PRESENT:

TITLE:

Items which do not conform are noted by:

Conformity it noted by a “√”

C- Critical K - Key O - Other NA- Not Applicable

Check the applicable analytical methods:

- | | |
|--------------------------|--|
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Seawater (APHA)[PART II] |
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Seawater using MA-1 [PART II] |
| <input type="checkbox"/> | Membrane Filtration Technique for Seawater using mTEC [PART II] |
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Shellfish Meats (APHA)[PART III] |
| <input type="checkbox"/> | Standard Plate Count for Shellfish Meats [PART III] |
| <input type="checkbox"/> | Elevated Temperature Coliform Plate Method for Shellfish Meats [PART III] |
| <input type="checkbox"/> | Male Specific Coliphage for Soft-shelled Clams and American Oysters [PART III] |

PART 1 - QUALITY ASSURANCE

CODE	REF.	ITEM	
K	8, 11	1.1 Quality Assurance (QA) Plan	
		<input type="checkbox"/>	1.1.1 Written Plan (Check those items which apply.)
		<input type="checkbox"/>	a. Organization of the laboratory.
		<input type="checkbox"/>	b. Staff training requirements.
		<input type="checkbox"/>	c. Standard operating procedures.
		<input type="checkbox"/>	d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		<input type="checkbox"/>	e. Laboratory safety.
		<input type="checkbox"/>	f. Internal performance assessment.
C	8	<input type="checkbox"/>	1.1.2 QA Plan Implemented.
K	11	<input type="checkbox"/>	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s) _____
1.2 Educational/Experience Requirements			
C	State's Human Resources Department	<input type="checkbox"/>	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	<input type="checkbox"/>	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three months of experience in laboratory sciences.
1.3 Work Area			
O	8,11	<input type="checkbox"/>	1.3.1 Adequate for workload and storage.
K	11	<input type="checkbox"/>	1.3.2 Clean, well-lighted.
K	11	<input type="checkbox"/>	1.3.3 Adequate temperature control.
O	11	<input type="checkbox"/>	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	<input type="checkbox"/>	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
1.4 Laboratory Equipment			
O	9	<input type="checkbox"/>	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	<input type="checkbox"/>	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	<input type="checkbox"/>	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	<input type="checkbox"/>	1.4.4 pH meter is calibrated daily or with each use Results are recorded and records maintained.
K	11	<input type="checkbox"/>	1.4.5 A minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	8,15	<input type="checkbox"/>	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt

				procedure or through determination of the slope. <i>(Circle the method used.)</i>
K	9	<input type="checkbox"/>	1.4.7	Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13	<input type="checkbox"/>	1.4.8	Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11	<input type="checkbox"/>	1.4.9	Refrigerator temperature(s) are monitored at least once daily on workdays. Results are recorded and records maintained.
K	1	<input type="checkbox"/>	1.4.10	Refrigerator temperature is maintained at 0 to 4°C.
C	9	<input type="checkbox"/>	1.4.11	The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11	<input type="checkbox"/>	1.4.12	Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9	<input type="checkbox"/>	1.4.13	Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11	<input type="checkbox"/>	1.4.14	Temperature of the waterbath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9	<input type="checkbox"/>	1.4.15	The thermometers used in the waterbath are graduated in at least 0.1°C increments.
C	13	<input type="checkbox"/>	1.4.16	The waterbath has adequate capacity for workload.
K	9	<input type="checkbox"/>	1.4.17	The level of water in the waterbath covers the level of liquid in the incubating tubes.
K	8, 11	<input type="checkbox"/>	1.4.18	Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4	<input type="checkbox"/>	1.4.19	All working thermometers are appropriately immersed.
C	29	<input type="checkbox"/>	1.4.20	Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	11	<input type="checkbox"/>	1.4.21	A mercury-in-glass standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9	<input type="checkbox"/>	1.4.22	Standards thermometers are checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination_____.
C	29	<input type="checkbox"/>	1.4.23	Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of ≤ ±0.05°C are used as the laboratory standards thermometer. <i>(Circle the thermometer type used.)</i>
K	13, 33	<input type="checkbox"/>	1.4.24	Incubator and water bath working thermometers are checked <u>verified</u> annually against the standards thermometer at the temperatures at which they are used. <u>Discard working temperature-sensing devices that differ by >1°C from the reference/standards device.</u> Results are recorded and records maintained.
O	11	<input type="checkbox"/>	1.4.25	Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing				
O	9	<input type="checkbox"/>	1.5.1	Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding materials.
K	9	<input type="checkbox"/>	1.5.2	Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.
K	9	<input type="checkbox"/>	1.5.3	Sample containers are made of glass or some other inert material.
O	9	<input type="checkbox"/>	1.5.4	Dilution bottles and tubes are made of borosilicate glass or plastic and closed

				with rubber stoppers, caps or screw caps with nontoxic liners.
K	9	<input type="checkbox"/>	1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9	<input type="checkbox"/>	1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1mL used to deliver 0.1 mL aliquots.
K	9	<input type="checkbox"/>	1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9	<input type="checkbox"/>	1.5.8	In washing reusable pipettes, a succession of at least three fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2	<input type="checkbox"/>	1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11	<input type="checkbox"/>	1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination				
K	9	<input type="checkbox"/>	1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8	<input type="checkbox"/>	1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	11, 30	<input type="checkbox"/>	1.6.3	The autoclave provides a sterilizing temperature of 121± 2°C as determined for each load using a calibrated maximum registering thermometer. As an alternative, an appropriate temperature monitoring device is used in place of the maximum registering thermometer when these are unavailable due to the ban on mercury.
K	11	<input type="checkbox"/>	1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16	<input type="checkbox"/>	1.6.5	The autoclave standards thermometer is checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory; or, is checked in-house at the steam point (100°C) if it has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination _____
K	1	<input type="checkbox"/>	1.6.6	Working autoclave thermometers are checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11	<input type="checkbox"/>	1.6.7	Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11	<input type="checkbox"/>	1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13	<input type="checkbox"/>	1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11	<input type="checkbox"/>	1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180°C.
K	9	<input type="checkbox"/>	1.6.11	A thermometer capable of determining temperatures accurately in the range of 160 to 180°C is used to monitor the operation of the hot-air sterilizing oven
K	13	<input type="checkbox"/>	1.6.12	Records of temperatures and exposure times are maintained for the operation of the hot-air sterilizing oven during use.
K	11	<input type="checkbox"/>	1.6.13	Spore strips/suspensions are used quarterly to evaluate the effectiveness of the

			sterilization process in the hot-air oven. Records are maintained.
K	11	<input type="checkbox"/>	1.6.14 Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C.
C	1	<input type="checkbox"/>	1.6.15 The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1	<input type="checkbox"/>	1.6.16 The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9	<input type="checkbox"/>	1.6.17 Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9	<input type="checkbox"/>	1.6.18 Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2 hours.
C	2	<input type="checkbox"/>	1.6.19 The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2	<input type="checkbox"/>	1.6.20 The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18	<input type="checkbox"/>	1.6.21 Hardwood applicator transfer sticks are properly sterilized. Method of sterilization _____
C	2	<input type="checkbox"/>	1.6.22 The sterility of the hardwood applicator transfer sticks is checked routinely. Results are recorded and the records maintained.
O	13	<input type="checkbox"/>	1.6.23 Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation			
K	3, 5	<input type="checkbox"/>	1.7.1 Media is commercially dehydrated except in the case of medium A-1 which must be prepared from the individual components and modified MacConkey agar which may be prepared from its components.
K	11	<input type="checkbox"/>	1.7.2 Media is prepared according to manufacturer's instructions.
O	11	<input type="checkbox"/>	1.7.3 Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11	<input type="checkbox"/>	1.7.4 Dehydrated media are labeled with date of receipt and date opened.
C	12	<input type="checkbox"/>	1.7.5 Caked or expired media or media components are discarded.
C	11	<input type="checkbox"/>	1.7.6 Reagent water is distilled or deionized (circle appropriate choice), tested monthly and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) or is less than 2.0 µSiemens/cm conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11	<input type="checkbox"/>	1.7.7 Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level (< 0.1 mg/L). Results are recorded and the records maintained. Specify method of determination _____.
K	11	<input type="checkbox"/>	1.7.8 Reagent water contains <100 CFU/mL as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11	<input type="checkbox"/>	1.7.9 Media prepared from commercial dehydrated components are sterilized according to the manufacturer's instructions.
K	9	<input type="checkbox"/>	1.7.10 The volume and concentration of media in the tube are suitable for the amount of sample inoculated.
C	11	<input type="checkbox"/>	1.7.11 Total time of exposure of sugar broths to autoclave temperatures does not exceed 45 minutes.
C	1	<input type="checkbox"/>	1.7.12 Media sterility is determined for each load sterilized. Results are recorded and the records maintained.
C	1	<input type="checkbox"/>	1.7.13 Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is

			made from its individual components.
O	9	<input type="checkbox"/>	1.7.14 Sterile phosphate buffered dilution water is used as the sample diluent.
K	11	<input type="checkbox"/>	1.7.15 The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media			
K	9	<input type="checkbox"/>	1.8.1 Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11	<input type="checkbox"/>	1.8.2 Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13	<input type="checkbox"/>	1.8.3 Stored media are labeled with the storage expiration date or the sterilization date.
K	9	<input type="checkbox"/>	1.8.4 Storage of prepared culture media at room temperature does not exceed 7 days.
K	2	<input type="checkbox"/>	1.8.5 Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed 1 month.
K	11	<input type="checkbox"/>	1.8.6 Storage under refrigeration of prepared culture media with screw-cap closures does not exceed 3 months.
K	17	<input type="checkbox"/>	1.8.7 All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES			
2.1 Collection and Transportation of Samples			
C	11	<input type="checkbox"/>	2.1.1 Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1	<input type="checkbox"/>	2.1.2 Samples are identified with collectors name, harvest area, sampling station, time and date of collection.
C	9	<input type="checkbox"/>	2.1.3 Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1	<input type="checkbox"/>	2.1.4 A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9	<input type="checkbox"/>	2.1.5 Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.
2.2 Bacteriological Examination of Seawater by the APHA MPN			
C	9	<input type="checkbox"/>	2.2.1 Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
C	2	<input type="checkbox"/>	2.2.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	9	<input type="checkbox"/>	2.2.3 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9	<input type="checkbox"/>	2.2.4 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	<input type="checkbox"/>	2.2.5 In a single dilution series not less than 12 tubes are used (for depuration at least 5 tubes are used).
C	6	<input type="checkbox"/>	2.2.6 In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____

			Range of MPN _____
			Strength of media used _____
K	9	<input type="checkbox"/>	2.2.7 Inoculated tubes are incubated in air at 35 ± 0.5°C.
C	2	<input type="checkbox"/>	2.2.8 Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained.
			Positive process control _____ Negative process control _____
K	9	<input type="checkbox"/>	2.2.9 Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
			2.3 Confirmed Test for Seawater by APHA MPN
C	9	<input type="checkbox"/>	2.3.1 Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9	<input type="checkbox"/>	2.3.2 EC medium is used as the confirmatory medium for fecal coliforms.
C	2	<input type="checkbox"/>	2.3.3 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
			Positive productivity control _____ Negative productivity control _____
K	9, 11	<input type="checkbox"/>	2.3.4 Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. <i>(Circle the method of transfer.)</i>
C	9	<input type="checkbox"/>	2.3.5 BGB tubes are incubated at 35 ± 0.5°C.
K	9	<input type="checkbox"/>	2.3.6 BGB tubes are read after 48 ± 3 hours of incubation.
C	9	<input type="checkbox"/>	2.3.7 EC tubes are incubated in a circulating waterbath maintained at 44.5 ± 0.2°C.
C	9	<input type="checkbox"/>	2.3.8 EC tubes are read after 24 ± 2 hours of incubation.
C	9	<input type="checkbox"/>	2.3.9 The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
			2.4 Computation of Results – APHA MPN
K	9	<input type="checkbox"/>	2.4.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7	<input type="checkbox"/>	2.4.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	<input type="checkbox"/>	2.4.3 Results are reported as MPN/100 mL of sample.
			2.5 Bacteriological Examination of Seawater by the MA-1 Method
C	5	<input type="checkbox"/>	2.5.1 A-1 medium complete is used in the analysis.
C	2, 31	<input type="checkbox"/>	2.5.2 A-1 medium without salicin is used in the analysis. Comparability testing supports use of A-1 medium without salicin. Study records are available.
C	5	<input type="checkbox"/>	2.5.3 A-1 medium sterilized for 10 minutes at 121°C.
C	2	<input type="checkbox"/>	2.5.4 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
			Positive productivity control _____ Negative productivity control _____
C	9	<input type="checkbox"/>	2.5.5 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9	<input type="checkbox"/>	2.5.6 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	<input type="checkbox"/>	2.5.7 In a single dilution series at least 12 tubes are used.

C	6	<input type="checkbox"/>	2.5.8	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	<input type="checkbox"/>	2.5.9	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and waterbath incubation Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	<input type="checkbox"/>	2.5.10	Inoculated tubes are placed in an air incubator at 35 ± 0.5°C for 3 ± 0.5 hours of resuscitation.
C	5	<input type="checkbox"/>	2.5.11	After 3 ± 0.5 hours resuscitation at 35°C, inoculated tubes are incubated at 44.5 ± 0.2°C in a circulating waterbath for the remainder of the 24 ± 2 hours.
C	5	<input type="checkbox"/>	2.5.12	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.6 Computation of Results – APHA MPN				
K	9	<input type="checkbox"/>	2.6.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	<input type="checkbox"/>	2.6.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	<input type="checkbox"/>	2.6.3	Results are reported as MPN/100 mL of sample.
2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment				
C	23, 24	<input type="checkbox"/>	2.7.1	When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at 44.5 ± 0.5°C under any loading capacity.
C	23	<input type="checkbox"/>	2.7.2	When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	<input type="checkbox"/>	2.7.3	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	<input type="checkbox"/>	2.7.4	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	<input type="checkbox"/>	2.7.5	Colonies are counted with the aid of magnification.
C	11, 23	<input type="checkbox"/>	2.7.6	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of 0.45 µm and certified by the manufacturer for fecal coliform analyses.
C	2	<input type="checkbox"/>	2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
C	2	<input type="checkbox"/>	2.7.8	When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison testing implemented. The results are recorded and this record is maintained.
K	2, 11	<input type="checkbox"/>	2.7.9	New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2	<input type="checkbox"/>	2.7.10	The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2	<input type="checkbox"/>	2.7.11	Membrane filters which are beyond their expiration date are not used.
O	11	<input type="checkbox"/>	2.7.12	Forceps tips are clean.
O	11	<input type="checkbox"/>	2.7.13	Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.

K	11	<input type="checkbox"/>	2.7.14	Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11	<input type="checkbox"/>	2.7.15	If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11	<input type="checkbox"/>	2.7.16	Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11	<input type="checkbox"/>	2.7.17	Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26	<input type="checkbox"/>	2.7.18	A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11	<input type="checkbox"/>	2.7.19	The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2	<input type="checkbox"/>	2.7.20	Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
2.8 Media Preparation and Storage – MF using mTEC Agar				
K	11	<input type="checkbox"/>	2.8.1	Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
C	11	<input type="checkbox"/>	2.8.2	The phosphate buffered saline is properly sterilized.
K	23	<input type="checkbox"/>	2.8.3	A sufficient amount of medium (4-5 mL) is used in each plate.
O	11	<input type="checkbox"/>	2.8.4	Refrigerated prepared plates are stored for no more than 2 weeks in sealed plastic bags or containers to minimize evaporation.
2.9 Sample Analyses - MF using mTEC Agar				
C	24	<input type="checkbox"/>	2.9.1	mTEC agar is used.
C	2	<input type="checkbox"/>	2.9.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	23	<input type="checkbox"/>	2.9.3	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration.
C	23	<input type="checkbox"/>	2.9.4	The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25	<input type="checkbox"/>	2.9.5	Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23	<input type="checkbox"/>	2.9.6	Sample volumes are filtered under vacuum.
K	26	<input type="checkbox"/>	2.9.7	The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26	<input type="checkbox"/>	2.9.8	The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.
C	23	<input type="checkbox"/>	2.9.9	The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11	<input type="checkbox"/>	2.9.10	Blanks are run at the beginning of filtration, after every 10th aliquot and at the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).
C	2, 11	<input type="checkbox"/>	2.9.11	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24	<input type="checkbox"/>	2.9.12	Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at 35 + 0.5°C for 2 hours of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5°C for 24 ± 2 hours.

C	11, 23, 24	<input type="checkbox"/>	2.9.13	After 2 hours of resuscitation at 35°C, the watertight, tightly sealed containers are transferred to a circulating waterbath at 44.5 + 0.2°C, submerged completely and incubated for 22-24 hours.
2.10 Computation of Results - MF using mTEC Agar				
C	23	<input type="checkbox"/>	2.10.1	All yellow, yellow-green or yellow-brown colonies are counted.
C	23	<input type="checkbox"/>	2.10.2	Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as >80 x 100/the volume of sample filtered.
C	2, 11, 23	<input type="checkbox"/>	2.10.3	When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11	<input type="checkbox"/>	2.10.4	The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11	<input type="checkbox"/>	2.10.5	Results are reported as CFU/100 mL of sample.
PART III - SHELLFISH SAMPLES				
3.1 Collection and Transportation of Samples				
C	9	<input type="checkbox"/>	3.1.1	A representative sample of shellstock is collected.
K	9	<input type="checkbox"/>	3.1.2	Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9	<input type="checkbox"/>	3.1.3	Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9	<input type="checkbox"/>	3.1.4	Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1	<input type="checkbox"/>	3.1.5	Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
3.2 Preparation of Shellfish for Examination				
K	2,11	<input type="checkbox"/>	3.2.1	Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2	<input type="checkbox"/>	3.2.2	Blades of shucking knives are not corroded.
O	9	<input type="checkbox"/>	3.2.3	The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	<input type="checkbox"/>	3.2.4	The faucet used for rinsing the shellstock does not contain an aerator.
K	9	<input type="checkbox"/>	3.2.5	Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	<input type="checkbox"/>	3.2.6	Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	9	<input type="checkbox"/>	3.2.7	Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9	<input type="checkbox"/>	3.2.8	Shellstock are not shucked directly through the hinge.
C	9	<input type="checkbox"/>	3.2.9	Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	<input type="checkbox"/>	3.2.10	At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.
K	9	<input type="checkbox"/>	3.2.11	A representative sample of at least 12 shellfish is used for the analysis.
K	2	<input type="checkbox"/>	3.2.12	The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
O	9	<input type="checkbox"/>	3.2.13	Sterile phosphate buffered dilution water is used as the sample diluent.

C	9	<input type="checkbox"/>	3.2.14	Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9	<input type="checkbox"/>	3.2.15	APHA <i>Recommended Procedures for the Examination of Sea Water And Shellfish</i> , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA				
C	9	<input type="checkbox"/>	3.3.1	Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. <i>(Circle the medium used.)</i>
C	2	<input type="checkbox"/>	3.3.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9	<input type="checkbox"/>	3.3.3	Immediately (within 2 minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9	<input type="checkbox"/>	3.3.4	No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
C	9	<input type="checkbox"/>	3.3.5	Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6	<input type="checkbox"/>	3.3.6	In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	<input type="checkbox"/>	3.3.7	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9	<input type="checkbox"/>	3.3.8	Inoculated media are incubated at $35 \pm 0.5^{\circ}\text{C}$.
K	10	<input type="checkbox"/>	3.3.9	Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA				
C	9	<input type="checkbox"/>	3.4.1	EC medium is used as the confirmatory medium.
C	2	<input type="checkbox"/>	3.4.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11	<input type="checkbox"/>	3.4.3	Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. <i>(Circle the method of transfer.)</i>
C	9	<input type="checkbox"/>	3.4.4	EC tubes are incubated in a circulating waterbath at $44.5 \pm 0.2^{\circ}\text{C}$
K	9	<input type="checkbox"/>	3.4.5	EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9	<input type="checkbox"/>	3.4.6	The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.
3.5 Computation of Results for MPN Analyses				
K	9	<input type="checkbox"/>	3.5.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.
K	7	<input type="checkbox"/>	3.5.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9	<input type="checkbox"/>	3.5.3	Results are reported as MPN/100 grams of sample.

3.6 Standard Plate Count Method			
O	20	<input type="checkbox"/>	3.6.1 A standard plate count (SPC) analysis may be performed in conjunction with the analysis for fecal coliform organisms.
K	9	<input type="checkbox"/>	3.6.2 In the standard plate count procedure at least four plates are used, duplicates of two dilutions. One of the dilutions should produce colonies of 30 to 300 per plate.
K	2	<input type="checkbox"/>	3.6.3 Fifteen to 20 mL of tempered sterile plate count agar is used per plate.
C	9	<input type="checkbox"/>	3.6.4 Agar tempering bath maintains the agar at 44-46°C.
C	9	<input type="checkbox"/>	3.6.5 An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9	<input type="checkbox"/>	3.6.6 Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in 7 seconds) before plating.
C	9	<input type="checkbox"/>	3.6.7 Not more than 1 mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11	<input type="checkbox"/>	3.6.8 Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9,21	<input type="checkbox"/>	3.6.9 Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four high.
K	9	<input type="checkbox"/>	3.6.10 Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1	<input type="checkbox"/>	3.6.11 A hand tally or its equivalent is used for accuracy in counting.
3.7 Computation of Results -SPC			
K	9	<input type="checkbox"/>	3.7.1 Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19	<input type="checkbox"/>	3.7.2 Colony counts are reported as CFU/g of sample.
3.8 Bacteriological Analysis of Shellfish Using the ETCP			
C	2,3	<input type="checkbox"/>	3.8.1 Prepared modified MacConkey agar is used on the day that it is made.
K	3	<input type="checkbox"/>	3.8.2 Double strength modified MacConkey agar is used.
C	3	<input type="checkbox"/>	3.8.3 Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3	<input type="checkbox"/>	3.8.4 Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50°C until used.
K	2, 3	<input type="checkbox"/>	3.8.5 Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3	<input type="checkbox"/>	3.8.6 The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9	<input type="checkbox"/>	3.8.7 The sample homogenate is cultured within 2 minutes of blending.
C	2,3	<input type="checkbox"/>	3.8.8 Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.
K	3	<input type="checkbox"/>	3.8.9 Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2,3, 22	<input type="checkbox"/>	3.8.10 The container is gently swirled or slowly inverted once to mix the contents, which are subsequently distributed uniformly over six plates.
C	1	<input type="checkbox"/>	3.8.11 Media and diluent sterility are determined with each use. Results are recorded and the records maintained.
C	1	<input type="checkbox"/>	3.8.12 Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture _____ Negative control culture _____
C	3, 13	<input type="checkbox"/>	3.8.13 When solidified, the plates are placed inverted into an air incubator at 45.5 ± 0.5°C for 18 to 30 hours of incubation.
C	2	<input type="checkbox"/>	3.8.14 Plates are stacked no more than three high in the incubator.

C	2	<input type="checkbox"/>	3.8.15	Appropriately diluted pour plated process control cultures <i>accompany each set of samples throughout incubation</i> . The results are recorded and the records maintained. Positive process control _____ Negative process control _____
3.9 Computation of Results - ETCP				
K	11	<input type="checkbox"/>	3.9.1	Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1	<input type="checkbox"/>	3.9.2	A hand tally or its equivalent is used to aid in counting.
C	3, 6	<input type="checkbox"/>	3.9.3	All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3	<input type="checkbox"/>	3.9.4	Results are reported as CFU/100 grams of sample.
Bacteriological Examination of Soft-shelled Clams and American Oysters for Male Specific Coliphage (MSC)				
3.10 MSC Equipment and Supplies				
K	30	<input type="checkbox"/>	3.10.1	Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold 100 – 125 mL.
C	27, 28	<input type="checkbox"/>	3.10.2	The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	9	<input type="checkbox"/>	3.10.3	The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
C	27, 28	<input type="checkbox"/>	3.10.4	Sterile 0.22 µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1	<input type="checkbox"/>	3.10.5	The sterility of each lot of pre-sterilized syringes and syringe filters is determined. Results are recorded and records maintained.
K	1	<input type="checkbox"/>	3.10.6	The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28	<input type="checkbox"/>	3.10.7	The balance used provides a sensitivity of at least mg (0.01g).
C	27, 28	<input type="checkbox"/>	3.10.8	The temperature of the incubator used is maintained at 36 ± 1°C.
C	28	<input type="checkbox"/>	3.10.9	Sterile disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
3.11 MSC Media Preparation				
K	28	<input type="checkbox"/>	3.11.1	Media preparation and sterilization is according to the validated method.
K	27, 28	<input type="checkbox"/>	3.11.2	Bottom agar, double strength soft agar and growth broth are prepared from their individual components.
K	27, 28	<input type="checkbox"/>	3.11.3	Soft agar is prepared double strength in volumes of 2.5 mL.
C	27, 28	<input type="checkbox"/>	3.11.4	The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
O	27, 28	<input type="checkbox"/>	3.11.5	Storage of the bottom agar under refrigeration does not exceed 1 month.
K	27, 28	<input type="checkbox"/>	3.11.6	Unsterilized soft agar is stored at -20 °C -15C for up to 3 months.
K	27, 28	<input type="checkbox"/>	3.11.7	The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.
K	27, 28	<input type="checkbox"/>	3.11.8	Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	27, 28	<input type="checkbox"/>	3.11.9	Bottom agar plates are allowed to reach room temperature before use.
3.12 Preparation of the Soft-Shelled Clams and American Oysters for MSC Analysis				
K	2,11	<input type="checkbox"/>	3.12.1	Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2	<input type="checkbox"/>	3.12.2	The blades of shucking knives are not corroded.
O	9	<input type="checkbox"/>	3.12.3	The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	<input type="checkbox"/>	3.12.4	The faucet used for rinsing the shellfish does not contain an aerator.

K	9	<input type="checkbox"/>	3.12.5	The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	<input type="checkbox"/>	3.12.6	The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9	<input type="checkbox"/>	3.12.7	Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9	<input type="checkbox"/>	3.12.8	Shellfish are not shucked through the hinge.
C	9	<input type="checkbox"/>	3.12.9	The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	<input type="checkbox"/>	3.12.10	A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19	<input type="checkbox"/>	3.12.11	The sample is weighed to the nearest 0.1 gram.
3.13 MSC Sample Analysis				
C	28	<input type="checkbox"/>	3.13.1	<i>E.coli Famp ATCC 700891</i> is the bacterial host strain used in this procedure.
K	27, 28	<input type="checkbox"/>	3.13.2	Host cell growth broth is tempered at $36 \pm 1^\circ\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	27, 28	<input type="checkbox"/>	3.13.3	Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^\circ\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	27, 28	<input type="checkbox"/>	3.13.4	After inoculation, the host cell growth broth culture is not shaken.
C	28	<input type="checkbox"/>	3.13.5	A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.
C	28	<input type="checkbox"/>	3.13.6	The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28	<input type="checkbox"/>	3.13.7	The elution mixture is homogenized at high speed for 180 seconds.
C	28	<input type="checkbox"/>	3.13.8	Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
C	28	<input type="checkbox"/>	3.13.9	The homogenized elution mixture is centrifuged for 15 minutes at 9000 x g at 4°C.
C	27, 28	<input type="checkbox"/>	3.13.10	The supernatant is pipetted off, weighed and the weight recorded.
C	27, 28	<input type="checkbox"/>	3.13.11	The supernatant is allowed to warm to room temperature about 20 to 30 minutes.
K	27, 28	<input type="checkbox"/>	3.13.12	The autoclaved soft agar is tempered and held at $51 \pm 1^\circ\text{C}$ throughout the period of sample analysis.
K	27, 28	<input type="checkbox"/>	3.13.13	Two hundred microliters (0.2 mL) of log phase host strain <i>E coli</i> is added to the tempering soft agar immediately prior to adding the sample supernatant.
K	27, 28	<input type="checkbox"/>	3.13.14	The sample supernatant is shaken or vortexed before being added to the tempering soft agar.
C	27, 28	<input type="checkbox"/>	3.13.15	2.5 mL of sample supernatant is added to each tube of tempering soft agar.
C	27, 28	<input type="checkbox"/>	3.13.16	The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28	<input type="checkbox"/>	3.13.17	The soft agar/sample supernatant/host cell mixture is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.
C	28	<input type="checkbox"/>	3.13.18	Ten (10) plates are used, 2.5 mL per plate for a total of 25 mL of supernatant analyzed per sample.
K	27, 28	<input type="checkbox"/>	3.13.19	Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are recorded and records maintained. Positive control _____
K	27, 28	<input type="checkbox"/>	3.13.20	Growth broth is used as the negative control or blank.
K	27, 28	<input type="checkbox"/>	3.13.21	Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2	<input type="checkbox"/>	3.13.22	A negative control plate is plated at the beginning and end of each set of samples analyzed.

K	27, 28	<input type="checkbox"/>	3.13.23 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28	<input type="checkbox"/>	3.13.24 All plates are incubated at 36 ± 1°C for 18 ± 2 hours.
3.14 Computation of Results - MSC			
C	27	<input type="checkbox"/>	3.14.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 32	<input type="checkbox"/>	3.14.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten plates, the count is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/ 100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as > 20,000 PFU/100 grams.
K	28	<input type="checkbox"/>	3.14.3 The formula used for determining the density of MSC in PFU/100 grams is: (0.364) (N) (Ws), where N = total number of plaques counted on all 10 plates and Ws = weight of the supernatant used.
O	9	<input type="checkbox"/>	3.14.4 The MSC count is rounded off conventionally to give a whole number.

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LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
MICROBIOLOGICAL COMPONENT: (Part I-III)	
A. Results	
Total # of Critical (C) Nonconformities in Parts I-III	_____
Total # of Key (K) Nonconformities in Parts I-III	_____
Total # of Critical, Key and Other (O)	_____
Nonconformities in Parts I-III	
B. Criteria for Determining Laboratory Status of the Microbiological Component:	
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <p style="margin-left: 40px;">a. The total # of Critical nonconformities is ≥ 4 or</p> <p style="margin-left: 40px;">b. The total # of Key nonconformities is ≥ 13 or</p> <p style="margin-left: 40px;">c. The total # of Critical, Key and Other is ≥ 18</p> <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but ≤ 3.</p>	
C. Laboratory Status (<i>circle appropriate</i>)	
Does Not Conform	Provisionally Conforms
Conforms	
Acknowledgment by Laboratory Director/Supervisor:	
All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____.	
Laboratory Signature: _____	Date: _____
LEO Signature: _____	Date: _____

NSSP Form LAB-100 Microbiology Rev. October 2015

**PUBLIC HEALTH SERVICE
U.S. FOOD AND DRUG ADMINISTRATION
OFFICE OF FOOD SAFETY
SHELLFISH AND AQUACULTURE POLICY BRANCH
5100 PAINT BRANCH PARKWAY
COLLEGE PARK, MD 20740-3835
TEL. 240- 402-2151/2055/4960 FAX 301-436-2601**

SHELLFISH LABORATORY EVALUATION CHECKLIST

LABORATORY:

ADDRESS:

TELEPHONE:

FAX:

EMAIL:

DATE OF EVALUATION:

DATE OF REPORT:

LAST EVALUATION:

LABORATORY REPRESENTED BY:

TITLE:

LABORATORY EVALUATION OFFICER:

SHELLFISH SPECIALIST:

OTHER OFFICIALS PRESENT:

REGION:

TITLE:

Items which do not conform are noted by:

Conformity it noted by a “√”

C- Critical K - Key O - Other NA- Not Applicable

Check the applicable analytical methods:

<input type="checkbox"/>	Multiple Tube Fermentation Technique for Seawater (APHA)[PART II]
<input type="checkbox"/>	Multiple Tube Fermentation Technique for Seawater using MA-1 [PART II]
<input type="checkbox"/>	Membrane Filtration Technique for Seawater using mTEC [PART II]
<input type="checkbox"/>	Multiple Tube Fermentation Technique for Shellfish Meats (APHA)[PART III]
<input checked="" type="checkbox"/>	Standard Plate Count for Shellfish Meats [PART III]
<input type="checkbox"/>	Elevated Temperature Coliform Plate Method for Shellfish Meats [PART III]
<input type="checkbox"/>	Male Specific Coliphage for Soft-shelled Clams and American Oysters [PART III]

PART 1 - QUALITY ASSURANCE

CODE	REF.	ITEM	
K	8, 11	1.1 Quality Assurance (QA) Plan	
		<input type="checkbox"/>	1.1.1 Written Plan (Check those items which apply.)
			a. Organization of the laboratory.
		<input type="checkbox"/>	b. Staff training requirements.
			c. Standard operating procedures.
		<input type="checkbox"/>	d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		<input type="checkbox"/>	e. Laboratory safety.
		<input type="checkbox"/>	f. Internal performance assessment.
			g. External performance assessment.
C	8	<input type="checkbox"/>	1.1.2 QA Plan Implemented.
K	11		1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s) _____
		1.2 Educational/Experience Requirements	
C	State's Human Resources Department	<input type="checkbox"/>	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	<input type="checkbox"/>	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	<input type="checkbox"/>	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three months of experience in laboratory sciences.
		1.3 Work Area	
O	8,11	<input type="checkbox"/>	1.3.1 Adequate for workload and storage.
K	11		1.3.2 Clean, well-lighted.
K	11	<input type="checkbox"/>	1.3.3 Adequate temperature control.
O	11	<input type="checkbox"/>	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	<input type="checkbox"/>	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
		1.4 Laboratory Equipment	
O	9	<input type="checkbox"/>	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	<input type="checkbox"/>	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	<input type="checkbox"/>	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	<input type="checkbox"/>	1.4.4 pH meter is calibrated daily or with each use Results are recorded and records maintained.
K	11	<input type="checkbox"/>	1.4.5 A minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	8,15	<input type="checkbox"/>	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt

			procedure or through determination of the slope. (<i>Circle the method used.</i>)
K	9	<input type="checkbox"/>	1.4.7 Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13	<input type="checkbox"/>	1.4.8 Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11	<input type="checkbox"/>	1.4.9 Refrigerator temperature(s) are monitored at least once daily on workdays Results are recorded and records maintained.
K	1	<input type="checkbox"/>	1.4.10 Refrigerator temperature is maintained at 0 to 4°C.
C	9	<input type="checkbox"/>	1.4.11 The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11	<input type="checkbox"/>	1.4.12 Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9	<input type="checkbox"/>	1.4.13 Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11	<input type="checkbox"/>	1.4.14 Temperature of the waterbath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9	<input type="checkbox"/>	1.4.15 The thermometers used in the waterbath are graduated in at least 0.1°C increments.
C	13	<input type="checkbox"/>	1.4.16 The waterbath has adequate capacity for workload.
K	9	<input type="checkbox"/>	1.4.17 The level of water in the waterbath covers the level of liquid in the incubating tubes.
K	8, 11	<input type="checkbox"/>	1.4.18 Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4	<input type="checkbox"/>	1.4.19 All working thermometers are appropriately immersed.
C	29, 33	<input type="checkbox"/>	1.4.20 Working thermometers are either: <u>calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers having the accuracy and tolerance of mercury,</u> or <u>appropriately calibrated low drift electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs) with an accuracy of less than or equal to ±0.05°C, with an accuracy and tolerance appropriate for the application.</u>
C	11	<input type="checkbox"/>	1.4.21 A mercury-in-glass standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9	<input type="checkbox"/>	1.4.22 Standards thermometers are checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination _____.
C	29	<input type="checkbox"/>	1.4.23 Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of ≤ ±0.05°C are used as the laboratory standards thermometer. (<i>Circle the thermometer type used.</i>)
K	13		1.4.24 Incubator and waterbath working thermometers are checked annually against the standards thermometer at the temperatures at which they are used. Results are recorded and records maintained.
O	11	<input type="checkbox"/>	1.4.25 Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing			
O	9	<input type="checkbox"/>	1.5.1 Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding materials.
K	9	<input type="checkbox"/>	1.5.2 Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.
K	9	<input type="checkbox"/>	1.5.3 Sample containers are made of glass or some other inert material.

O	9	<input type="checkbox"/>	1.5.4	Dilution bottles and tubes are made of borosilicate glass or plastic and closed with rubber stoppers, caps or screw caps with nontoxic liners.
K	9	<input type="checkbox"/>	1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9	<input type="checkbox"/>	1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1mL used to deliver 0.1 mL aliquots.
K	9	<input type="checkbox"/>	1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9	<input type="checkbox"/>	1.5.8	In washing reusable pipettes, a succession of at least three fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2	<input type="checkbox"/>	1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11		1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination				
K	9	<input type="checkbox"/>	1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8		1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	11, 30	<input type="checkbox"/>	1.6.3	The autoclave provides a sterilizing temperature of 121± 2°C as determined for each load using a calibrated maximum registering thermometer. As an alternative, an appropriate temperature monitoring device is used in place of the maximum registering thermometer when these are unavailable due to the ban on mercury.
K	11	<input type="checkbox"/>	1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16	<input type="checkbox"/>	1.6.5	The autoclave standards thermometer is checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory; or, is checked in-house at the steam point (100°C) if it has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination _____
K	1	<input type="checkbox"/>	1.6.6	Working autoclave thermometers are checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11	<input type="checkbox"/>	1.6.7	Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11		1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13	<input type="checkbox"/>	1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11		1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180°C.
K	9	<input type="checkbox"/>	1.6.11	A thermometer capable of determining temperatures accurately in the range of 160 to 180°C is used to monitor the operation of the hot-air sterilizing oven
K	13	<input type="checkbox"/>	1.6.12	Records of temperatures and exposure times are maintained for the operation of the hot-air sterilizing oven during use.

K	11	<input type="checkbox"/>	1.6.13 Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Records are maintained.
K	11	<input type="checkbox"/>	1.6.14 Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C.
C	1	<input type="checkbox"/>	1.6.15 The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1	<input type="checkbox"/>	1.6.16 The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9	<input type="checkbox"/>	1.6.17 Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9	<input type="checkbox"/>	1.6.18 Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2 hours.
C	2	<input type="checkbox"/>	1.6.19 The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2	<input type="checkbox"/>	1.6.20 The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18	<input type="checkbox"/>	1.6.21 Hardwood applicator transfer sticks are properly sterilized. Method of sterilization _____
C	2	<input type="checkbox"/>	1.6.22 The sterility of the hardwood applicator transfer sticks is checked routinely. Results are recorded and the records maintained.
O	13	<input type="checkbox"/>	1.6.23 Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation			
K	3, 5	<input type="checkbox"/>	1.7.1 Media is commercially dehydrated except in the case of medium A-1 which must be prepared from the individual components and modified MacConkey agar which may be prepared from its components.
K	11	<input type="checkbox"/>	1.7.2 Media is prepared according to manufacturer's instructions.
O	11	<input type="checkbox"/>	1.7.3 Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11	<input type="checkbox"/>	1.7.4 Dehydrated media are labeled with date of receipt and date opened.
C	12	<input type="checkbox"/>	1.7.5 Caked or expired media or media components are discarded.
C	11	<input type="checkbox"/>	1.7.6 Reagent water is distilled or deionized (circle appropriate choice), tested monthly and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) or is less than 2.0 µSiemens/cm conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11	<input type="checkbox"/>	1.7.7 Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level (< 0.1 mg/L). Results are recorded and the records maintained. Specify method of determination _____.
K	11	<input type="checkbox"/>	1.7.8 Reagent water contains <100 CFU/mL as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11	<input type="checkbox"/>	1.7.9 Media prepared from commercial dehydrated components are sterilized according to the manufacturer's instructions.
K	9	<input type="checkbox"/>	1.7.10 The volume and concentration of media in the tube are suitable for the amount of sample inoculated.
C	11	<input type="checkbox"/>	1.7.11 Total time of exposure of sugar broths to autoclave temperatures does not exceed 45 minutes.
C	1	<input type="checkbox"/>	1.7.12 Media sterility is determined for each load sterilized. Results are recorded and the records maintained.
C	1	<input type="checkbox"/>	1.7.13 Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated

			media received or with each batch of media prepared when the medium is made from its individual components.
O	9	<input type="checkbox"/>	1.7.14 Sterile phosphate buffered dilution water is used as the sample diluent.
K	11	<input type="checkbox"/>	1.7.15 The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media			
K	9	<input type="checkbox"/>	1.8.1 Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11	<input type="checkbox"/>	1.8.2 Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13	<input type="checkbox"/>	1.8.3 Stored media are labeled with the storage expiration date or the sterilization date.
K	9		1.8.4 Storage of prepared culture media at room temperature does not exceed 7 days.
K	2	<input type="checkbox"/>	1.8.5 Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed 1 month.
K	11	<input type="checkbox"/>	1.8.6 Storage under refrigeration of prepared culture media with screw-cap closures does not exceed 3 months.
K	17	<input type="checkbox"/>	1.8.7 All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES			
2.1 Collection and Transportation of Samples			
C	11	<input type="checkbox"/>	2.1.1 Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1	<input type="checkbox"/>	2.1.2 Samples are identified with collectors name, harvest area, sampling station, time and date of collection.
C	9	<input type="checkbox"/>	2.1.3 Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1		2.1.4 A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9	<input type="checkbox"/>	2.1.5 Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.
2.2 Bacteriological Examination of Seawater by the APHA MPN			
C	9	<input type="checkbox"/>	2.2.1 Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
C	2	<input type="checkbox"/>	2.2.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	9	<input type="checkbox"/>	2.2.3 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9	<input type="checkbox"/>	2.2.4 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	<input type="checkbox"/>	2.2.5 In a single dilution series not less than 12 tubes are used (for deputation at least 5 tubes are used).
C	6	<input type="checkbox"/>	2.2.6 In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____

			Range of MPN _____
			Strength of media used _____
K	9	<input type="checkbox"/>	2.2.7 Inoculated tubes are incubated in air at 35 ± 0.5°C.
C	2	<input type="checkbox"/>	2.2.8 Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained.
			Positive process control _____ Negative process control _____
K	9	<input type="checkbox"/>	2.2.9 Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
			2.3 Confirmed Test for Seawater by APHA MPN
C	9	<input type="checkbox"/>	2.3.1 Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9	<input type="checkbox"/>	2.3.2 EC medium is used as the confirmatory medium for fecal coliforms.
C	2	<input type="checkbox"/>	2.3.3 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
			Positive productivity control _____ Negative productivity control _____
K	9, 11	<input type="checkbox"/>	2.3.4 Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. <i>(Circle the method of transfer.)</i>
C	9	<input type="checkbox"/>	2.3.5 BGB tubes are incubated at 35 ± 0.5°C.
K	9	<input type="checkbox"/>	2.3.6 BGB tubes are read after 48 ± 3 hours of incubation.
C	9	<input type="checkbox"/>	2.3.7 EC tubes are incubated in a circulating waterbath maintained at 44.5 ± 0.2°C.
C	9	<input type="checkbox"/>	2.3.8 EC tubes are read after 24 ± 2 hours of incubation.
C	9	<input type="checkbox"/>	2.3.9 The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
			2.4 Computation of Results – APHA MPN
K	9	<input type="checkbox"/>	2.4.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7	<input type="checkbox"/>	2.4.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	<input type="checkbox"/>	2.4.3 Results are reported as MPN/100 mL of sample.
			2.5 Bacteriological Examination of Seawater by the MA-1 Method
C	5	<input type="checkbox"/>	2.5.1 A-1 medium complete is used in the analysis.
C	2, 31	<input type="checkbox"/>	2.5.2 A-1 medium without salicin is used in the analysis. Comparability testing supports use of A-1 medium without salicin. Study records are available.
C	5	<input type="checkbox"/>	2.5.3 A-1 medium sterilized for 10 minutes at 121°C.
C	2	<input type="checkbox"/>	2.5.4 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
			Positive productivity control _____ Negative productivity control _____
C	9	<input type="checkbox"/>	2.5.5 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9	<input type="checkbox"/>	2.5.6 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	<input type="checkbox"/>	2.5.7 In a single dilution series at least 12 tubes are used.

C	6	<input type="checkbox"/>	2.5.8 In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	<input type="checkbox"/>	2.5.9 Appropriately diluted process control cultures accompany the samples <i>throughout both resuscitation and waterbath incubation</i> Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	<input type="checkbox"/>	2.5.10 Inoculated tubes are placed in an air incubator at $35 \pm 0.5^\circ\text{C}$ for 3 ± 0.5 hours of resuscitation.
C	5	<input type="checkbox"/>	2.5.11 After 3 ± 0.5 hours resuscitation at 35°C , inoculated tubes are incubated at $44.5 \pm 0.2^\circ\text{C}$ in a circulating waterbath for the remainder of the 24 ± 2 hours.
C	5	<input type="checkbox"/>	2.5.12 The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.6 Computation of Results – APHA MPN			
K	9	<input type="checkbox"/>	2.6.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	<input type="checkbox"/>	2.6.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9		2.6.3 Results are reported as MPN/100 mL of sample.
2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment			
C	23, 24	<input type="checkbox"/>	2.7.1 When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at $44.5 \pm 0.5^\circ\text{C}$ under any loading capacity.
C	23	<input type="checkbox"/>	2.7.2 When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	<input type="checkbox"/>	2.7.3 Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	<input type="checkbox"/>	2.7.4 The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	<input type="checkbox"/>	2.7.5 Colonies are counted with the aid of magnification.
C	11, 23		2.7.6 Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of $0.45 \mu\text{m}$ and certified by the manufacturer for fecal coliform analyses.
C	2		2.7.7 Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
C	2	<input checked="" type="checkbox"/>	2.7.8 When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison testing implemented. The results are recorded and this record is maintained.
K	2, 11	<input type="checkbox"/>	2.7.9 New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2	<input type="checkbox"/>	2.7.10 The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2	<input type="checkbox"/>	2.7.11 Membrane filters which are beyond their expiration date are not used.
O	11		2.7.12 Forceps tips are clean.
O	11	<input type="checkbox"/>	2.7.13 Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.

K	11	<input type="checkbox"/>	2.7.14 Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11		2.7.15 If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11	<input type="checkbox"/>	2.7.16 Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11	<input type="checkbox"/>	2.7.17 Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26	<input type="checkbox"/>	2.7.18 A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11	<input type="checkbox"/>	2.7.19 The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2	<input type="checkbox"/>	2.7.20 Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
2.8 Media Preparation and Storage – MF using mTEC Agar			
K	11	<input type="checkbox"/>	2.8.1 Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
C	11		2.8.2 The phosphate buffered saline is properly sterilized.
K	23	<input type="checkbox"/>	2.8.3 A sufficient amount of medium (4-5 mL) is used in each plate.
O	11		2.8.4 Refrigerated prepared plates are stored for no more than 2 weeks in sealed plastic bags or containers to minimize evaporation.
2.9 Sample Analyses - MF using mTEC Agar			
C	24	<input type="checkbox"/>	2.9.1 mTEC agar is used.
C	2	<input type="checkbox"/>	2.9.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	23	<input type="checkbox"/>	2.9.3 The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration.
C	23	<input type="checkbox"/>	2.9.4 The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25		2.9.5 Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23	<input type="checkbox"/>	2.9.6 Sample volumes are filtered under vacuum.
K	26		2.9.7 The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26	<input type="checkbox"/>	2.9.8 The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.
C	23	<input type="checkbox"/>	2.9.9 The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11	<input type="checkbox"/>	2.9.10 Blanks are run at the beginning of filtration, after every 10 th aliquot and at the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).
C	2, 11	<input type="checkbox"/>	2.9.11 Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24	<input type="checkbox"/>	2.9.12 Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at 35 + 0.5°C for 2 hours of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5°C for 24 ± 2 hours.

C	11, 23, 24	<input type="checkbox"/>	2.9.13 After 2 hours of resuscitation at 35°C, the watertight, tightly sealed containers are transferred to a circulating waterbath at 44.5 + 0.2°C, submerged completely and incubated for 22-24 hours.
			2.10 Computation of Results - MF using mTEC Agar
C	23	<input type="checkbox"/>	2.10.1 All yellow, yellow-green or yellow-brown colonies are counted.
C	23	<input type="checkbox"/>	2.10.2 Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as >80 x 100/the volume of sample filtered.
C	2, 11, 23	<input type="checkbox"/>	2.10.3 When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11	<input type="checkbox"/>	2.10.4 The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11	<input type="checkbox"/>	2.10.5 Results are reported as CFU/100 mL of sample.
PART III - SHELLFISH SAMPLES			
3.1 Collection and Transportation of Samples			
C	9	<input type="checkbox"/>	3.1.1 A representative sample of shellstock is collected.
K	9	<input type="checkbox"/>	3.1.2 Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9	<input type="checkbox"/>	3.1.3 Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9	<input type="checkbox"/>	3.1.4 Immediately after collection, shellfish samples are placed in dry storage(ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1	<input type="checkbox"/>	3.1.5 Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
3.2 Preparation of Shellfish for Examination			
K	2,11	<input type="checkbox"/>	3.2.1 Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2	<input type="checkbox"/>	3.2.2 Blades of shucking knives are not corroded.
O	9	<input type="checkbox"/>	3.2.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	<input type="checkbox"/>	3.2.4 The faucet used for rinsing the shellstock does not contain an aerator.
K	9	<input type="checkbox"/>	3.2.5 Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	<input type="checkbox"/>	3.2.6 Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	9	<input type="checkbox"/>	3.2.7 Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9	<input type="checkbox"/>	3.2.8 Shellstock are not shucked directly through the hinge.
C	9	<input type="checkbox"/>	3.2.9 Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	<input type="checkbox"/>	3.2.10 At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.
K	9	<input type="checkbox"/>	3.2.11 A representative sample of at least 12 shellfish is used for the analysis.
K	2	<input type="checkbox"/>	3.2.12 The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
O	9	<input type="checkbox"/>	3.2.13 Sterile phosphate buffered dilution water is used as the sample diluent.

C	9	<input type="checkbox"/>	3.2.14 Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9		3.2.15 APHA <i>Recommended Procedures for the Examination of Sea Water And Shellfish</i> , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA			
C	9	<input type="checkbox"/>	3.3.1 Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. (Circle the medium used.)
C	2	<input type="checkbox"/>	3.3.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9	<input type="checkbox"/>	3.3.3 Immediately (within 2 minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9	<input type="checkbox"/>	3.3.4 No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
C	9	<input type="checkbox"/>	3.3.5 Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6	<input type="checkbox"/>	3.3.6 In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2		3.3.7 Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9	<input type="checkbox"/>	3.3.8 Inoculated media are incubated at $35 \pm 0.5^{\circ}\text{C}$.
K	10	<input type="checkbox"/>	3.3.9 Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA			
C	9	<input type="checkbox"/>	3.4.1 EC medium is used as the confirmatory medium.
C	2	<input type="checkbox"/>	3.4.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11	<input type="checkbox"/>	3.4.3 Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. (Circle the method of transfer.)
C	9	<input type="checkbox"/>	3.4.4 EC tubes are incubated in a circulating waterbath at $44.5 \pm 0.2^{\circ}\text{C}$
K	9		3.4.5 EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9	<input type="checkbox"/>	3.4.6 The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.
3.5 Computation of Results for MPN Analyses			
K	9	<input type="checkbox"/>	3.5.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.
K	7	<input type="checkbox"/>	3.5.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9	<input type="checkbox"/>	3.5.3 Results are reported as MPN/100 grams of sample.

3.6 Standard Plate Count Method			
O	20	<input type="checkbox"/>	3.6.1 A standard plate count (SPC) analysis may be performed in conjunction with the analysis for fecal coliform organisms.
K	9	<input type="checkbox"/>	3.6.2 In the standard plate count procedure at least four plates are used, duplicates of two dilutions. One of the dilutions should produce colonies of 30 to 300 per plate.
K	2	<input type="checkbox"/>	3.6.3 Fifteen to 20 mL of tempered sterile plate count agar is used per plate.
C	9		3.6.4 Agar tempering bath maintains the agar at 44-46°C.
C	9	<input type="checkbox"/>	3.6.5 An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9	<input type="checkbox"/>	3.6.6 Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in 7 seconds) before plating.
C	9	<input type="checkbox"/>	3.6.7 Not more than 1 mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11	<input type="checkbox"/>	3.6.8 Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9,21	<input type="checkbox"/>	3.6.9 Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four high.
K	9	<input type="checkbox"/>	3.6.10 Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1	<input type="checkbox"/>	3.6.11 A hand tally or its equivalent is used for accuracy in counting.
3.7 Computation of Results -SPC			
K	9	<input type="checkbox"/>	3.7.1 Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19		3.7.2 Colony counts are reported as CFU/g of sample.
3.8 Bacteriological Analysis of Shellfish Using the ETCP			
C	2,3	<input checked="" type="checkbox"/>	3.8.1 Prepared modified MacConkey agar is used on the day that it is made.
K	3		3.8.2 Double strength modified MacConkey agar is used.
C	3	<input type="checkbox"/>	3.8.3 Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3	<input type="checkbox"/>	3.8.4 Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50°C until used.
K	2, 3	<input type="checkbox"/>	3.8.5 Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3		3.8.6 The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9	<input type="checkbox"/>	3.8.7 The sample homogenate is cultured within 2 minutes of blending.
C	2,3		3.8.8 Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.
K	3	<input type="checkbox"/>	3.8.9 Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2,3, 22	<input type="checkbox"/>	3.8.10 The container is gently swirled or slowly inverted once to mix the contents, which are subsequently distributed uniformly over six plates.
C	1	<input type="checkbox"/>	3.8.11 Media and diluent sterility are determined with each use. Results are recorded and the records maintained.
C	1	<input type="checkbox"/>	3.8.12 Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture _____ Negative control culture _____
C	3, 13	<input type="checkbox"/>	3.8.13 When solidified, the plates are placed inverted into an air incubator at 45.5 ± 0.5°C for 18 to 30 hours of incubation.
C	2	<input type="checkbox"/>	3.8.14 Plates are stacked no more than three high in the incubator.

C	2	<input type="checkbox"/>	3.8.15 Appropriately diluted pour plated process control cultures <i>accompany each set of samples throughout incubation</i> . The results are recorded and the records maintained. Positive process control _____ Negative process control _____
3.9 Computation of Results - ETCP			
K	11	<input type="checkbox"/>	3.9.1 Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1	<input type="checkbox"/>	3.9.2 A hand tally or its equivalent is used to aid in counting.
C	3, 6	<input type="checkbox"/>	3.9.3 All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3	<input type="checkbox"/>	3.9.4 Results are reported as CFU/100 grams of sample.
Bacteriological Examination of Soft-shelled Clams and American Oysters for Male Specific Coliphage (MSC)			
3.10 MSC Equipment and Supplies			
K	30	<input type="checkbox"/>	3.10.1 Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold 100 – 125 mL.
C	27, 28	<input type="checkbox"/>	3.10.2 The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	9	<input type="checkbox"/>	3.10.3 The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
C	27, 28	<input type="checkbox"/>	3.10.4 Sterile 0.22 µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1	<input type="checkbox"/>	3.10.5 The sterility of each lot of pre-sterilized syringes and syringe filters is determined. Results are recorded and records maintained.
K	1	<input type="checkbox"/>	3.10.6 The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28	<input type="checkbox"/>	3.10.7 The balance used provides a sensitivity of at least mg (0.01g).
C	27, 28	<input type="checkbox"/>	3.10.8 The temperature of the incubator used is maintained at 36 ± 1°C.
C	28	<input type="checkbox"/>	3.10.9 Sterile disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
3.11 MSC Media Preparation			
K	28	<input type="checkbox"/>	3.11.1 Media preparation and sterilization is according to the validated method.
K	27, 28	<input type="checkbox"/>	3.11.2 Bottom agar, double strength soft agar and growth broth are prepared from their individual components.
K	27, 28	<input type="checkbox"/>	3.11.3 Soft agar is prepared double strength in volumes of 2.5 mL.
C	27, 28	<input type="checkbox"/>	3.11.4 The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
O	27, 28	<input type="checkbox"/>	3.11.5 Storage of the bottom agar under refrigeration does not exceed 1 month.
K	27, 28	<input type="checkbox"/>	3.11.6 Unsterilized soft agar is stored at -20 °C -15C for up to 3 months.
K	27, 28	<input type="checkbox"/>	3.11.7 The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.
K	27, 28	<input type="checkbox"/>	3.11.8 Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	27, 28	<input type="checkbox"/>	3.11.9 Bottom agar plates are allowed to reach room temperature before use.
3.12 Preparation of the Soft-Shell Clams and American Oysters for MSC Analysis			
K	2,11	<input type="checkbox"/>	3.12.1 Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2	<input type="checkbox"/>	3.12.2 The blades of shucking knives are not corroded.
O	9	<input type="checkbox"/>	3.12.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	<input type="checkbox"/>	3.12.4 The faucet used for rinsing the shellfish does not contain an aerator.

K	9	<input type="checkbox"/>	3.12.5 The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	<input type="checkbox"/>	3.12.6 The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9	<input type="checkbox"/>	3.12.7 Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9	<input type="checkbox"/>	3.12.8 Shellfish are not shucked through the hinge.
C	9	<input type="checkbox"/>	3.12.9 The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	<input type="checkbox"/>	3.12.10 A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19	<input type="checkbox"/>	3.12.11 The sample is weighed to the nearest 0.1 gram.
3.13 MSC Sample Analysis			
C	28	<input type="checkbox"/>	3.13.1 E.coli <i>Famp</i> ATCC 700891 is the bacterial host strain used in this procedure.
K	27, 28	<input type="checkbox"/>	3.13.2 Host cell growth broth is tempered at $36 \pm 1^\circ\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	27, 28	<input type="checkbox"/>	3.13.3 Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^\circ\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	27, 28	<input type="checkbox"/>	3.13.4 After inoculation, the host cell growth broth culture is not shaken.
C	28	<input type="checkbox"/>	3.13.5 A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.
C	28	<input type="checkbox"/>	3.13.6 The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28	<input type="checkbox"/>	3.13.7 The elution mixture is homogenized at high speed for 180 seconds.
C	28	<input type="checkbox"/>	3.13.8 Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
C	28	<input type="checkbox"/>	3.13.9 The homogenized elution mixture is centrifuged for 15 minutes at 9000 x g at 4°C.
C	27, 28	<input type="checkbox"/>	3.13.10 The supernatant is pipetted off, weighed and the weight recorded.
C	27, 28	<input type="checkbox"/>	3.13.11 The supernatant is allowed to warm to room temperature about 20 to 30 minutes.
K	27, 28	<input type="checkbox"/>	3.13.12 The autoclaved soft agar is tempered and held at $51 \pm 1^\circ\text{C}$ throughout the period of sample analysis.
K	27, 28	<input type="checkbox"/>	3.13.13 Two hundred microliters (0.2 mL) of log phase host strain <i>E coli</i> is added to the tempering soft agar immediately prior to adding the sample supernatant.
K	27, 28	<input type="checkbox"/>	3.13.14 The sample supernatant is shaken or vortexed before being added to the tempering soft agar.
C	27, 28	<input type="checkbox"/>	3.13.15 2.5 mL of sample supernatant is added to each tube of tempering soft agar.
C	27, 28	<input type="checkbox"/>	3.13.16 The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28	<input type="checkbox"/>	3.13.17 The soft agar/sample supernatant/host cell mixture is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.
C	28	<input type="checkbox"/>	3.13.18 Ten (10) plates are used, 2.5 mL per plate for a total of 25 mL of supernatant analyzed per sample.
K	27, 28	<input type="checkbox"/>	3.13.19 Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are recorded and records maintained. Positive control _____
K	27, 28	<input type="checkbox"/>	3.13.20 Growth broth is used as the negative control or blank.
K	27, 28	<input type="checkbox"/>	3.13.21 Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2	<input type="checkbox"/>	3.13.22 A negative control plate is plated at the beginning and end of each set of samples analyzed.

K	27, 28	<input type="checkbox"/>	3.13.23 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28	<input type="checkbox"/>	3.13.24 All plates are incubated at $36 \pm 1^\circ\text{C}$ for 18 ± 2 hours.
3.14 Computation of Results - MSC			
C	27	<input type="checkbox"/>	3.14.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 32	<input type="checkbox"/>	3.14.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten plates, the count is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as $> 20,000$ PFU/100 grams.
K	28		3.14.3 The formula used for determining the density of MSC in PFU/100 grams is: $(0.364) (N) (W_s)$, where N = total number of plaques counted on all 10 plates and W_s = weight of the supernatant used.
O	9	<input type="checkbox"/>	3.14.4 The MSC count is rounded off conventionally to give a whole number.

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- ~~32-33.~~ NIST Monograph 150 states “the accuracy attainable is principally limited by the characteristics of the thermometer itself.”

LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
MICROBIOLOGICAL COMPONENT: (Part I-III)	
A. Results	
Total # of Critical (C) Nonconformities in Parts I-III	_____
Total # of Key (K) Nonconformities in Parts I-III	_____
Total # of Critical, Key and Other (O)	_____
Nonconformities in Parts I-III	
B. Criteria for Determining Laboratory Status of the Microbiological Component:	
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <ul style="list-style-type: none"> a. The total # of Critical nonconformities is ≥ 4 or b. The total # of Key nonconformities is ≥ 13 or c. The total # of Critical, Key and Other is ≥ 18 <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but ≤ 3.</p>	
C. Laboratory Status (circle appropriate)	
Does Not Conform	Provisionally Conforms
Conforms	
Acknowledgment by Laboratory Director/Supervisor:	
All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____.	
Laboratory Signature: _____	Date: _____
LEO Signature: _____	Date: _____

NSSP Form LAB-100 Microbiology Rev. October 2015

PUBLIC HEALTH SERVICE U.S. FOOD AND DRUG ADMINISTRATION OFFICE OF FOOD SAFETY SHELLFISH AND AQUACULTURE POLICY BRANCH 5001 CAMPUS DRIVE COLLEGE PARK, MD 20740-3835 TEL. 240-402-2151/2055/4960 FAX 301-436-2601		
SHELLFISH LABORATORY EVALUATION CHECKLIST Diarrhetic Shellfish Poisoning Toxins (DSP) LC-MS/MS		
LABORATORY:		
ADDRESS:		
TELEPHONE:	FAX:	EMAIL:
DATE OF EVALUATION:	DATE OF REPORT:	LAST EVALUATION:
LABORATORY REPRESENTED BY:		TITLE:
LABORATORY EVALUATION OFFICER:		SHELLFISH SPECIALIST:
OTHER OFFICIALS PRESENT:		TITLE:
Items which do not conform are noted by: C – Critical K - Key O - Other NA - Not Applicable Conformity is noted by a “1”		

PART I – QUALITY ASSURANCE		
Code	REF	Item Description
1.1 Quality Assurance (QA) Plan		
K	1, 7, 8	1.1.1 Written Plan adequately covers all the following: (check those that apply) a. Organization of the laboratory b. Staff training requirements c. Standard operating procedures d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance and rejection criteria established e. Laboratory safety f. Internal performance assessment g. External performance assessment
C	5	1.1.2 QA Plan is implemented.
1.2 Educational/Experience Requirements		
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial/private laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, chemistry, or another appropriate discipline with at least two (2) years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial/private laboratories, the analyst must have at least a high school diploma and shall have at least three (3) months of experience in laboratory sciences.
C	3	1.2.5 LC-MS Operator must be trained in the operation and maintenance of the specific liquid chromatography-mass spectrometry system used.
1.3 Work Area		
O	1	1.3.1 Adequate for workload and storage.
O	1	1.3.2 Clean and well lighted.
O	1	1.3.3 Adequate temperature control.
O	8	1.3.4 All work surfaces are nonporous and easily cleaned.

1.4 Laboratory Equipment		
C	3	1.4.1 A heat block or water bath capable of heating samples to 76 ± 2 °C.
K	2	1.4.2 Balances provide an appropriate sensitivity at the weights of use, at least 0.1 g for laboratory precision balances and 0.1 mg for analytical balances.
K	7, 8	1.4.3 The balance calibration is checked monthly using NIST class S, ASTM class 1 or 2 weights or equivalent. Results are recorded and records are maintained.
K	1	1.4.4 Refrigerator temperature is maintained between 0 and 4 °C.
K	7	1.4.5 Refrigerator temperature is monitored at least once daily. Results are recorded and records maintained.
K	2	1.4.6 Freezer temperature is maintained at -10 °C or below.
K	7	1.4.7 Freezer temperature is monitored at least once daily. Results are recorded and records maintained.
C	10	1.4.8 All in-service thermometers are properly calibrated and immersed.
K	4	1.4.9 All glassware is clean.
K	3, 12	1.4.10 An ultra-performance liquid chromatography system (UPLC) equipped with the following is used: <ul style="list-style-type: none"> a. mobile phase system <u>capable of</u> delivering a pulse-free flow of 0.12 mL/min b. solvent degasser (<u>optional</u>) c. autosampler (refrigerated preferred) with loop suitable for five (5) µL injections d. column compartment capable of controlling temperature at 40 °C e. a data collection system (e.g., computer, integrator)
C	3	1.4.11 A mass spectrometer equipped with the following is used: <ul style="list-style-type: none"> a. an electrospray ionization source operating in negative ion mode and b. multiple reaction monitoring scan mode capability. c. if a divert valve is used to divert LC flow at the beginning and end of each chromatographic run, the switching time should be at least one minute before the first peak elution and at least one minute after the last peak elution.
K	2	1.4.12 Autopipettors are calibrated for the appropriate volumes used and checked annually for accuracy. Results are recorded and records are maintained.
K	3	1.4.13 A centrifuge capable of generating 2000 x g and holding 15 mL and 50 mL polypropylene tubes is used.

		1.5 Reagents and Reference Solution Preparation and Storage
C	3	1.5.1 All solvents and reagents used are analytical or LC grade materials.
O	7	1.5.2 Water contains < 100 CFU/ml determined monthly using the heterotrophic plate count method. Results are recorded and records are maintained. (Not required for bottled reagent grade or HPLC grade water when used immediately upon opening. If the bottle of water is not used entirely immediately, the water must be tested as above prior to continued use.)
K	7	1.5.3 Reagents are properly stored and labeled with the date of receipt, date opened or date prepared and expiration date.
C	3	1.5.4 The mobile phase system used to analyze DSP toxins consists of: A: 2 mM ammonium formate and 50 mM formic acid in water B: 2 mM ammonium formate and 50 mM formic acid in 95% acetonitrile/5% water
O	2	1.5.5 Mobile phase is filtered <u>degassed manually</u> before use if the UPLC does not have a degasser <u>or if the degasser is not in use.</u>
C	3	1.5.6 Only certified reference materials are used for standard solutions. Source of the reference standard: _____
C	6	1.5.7 All primary standards are stored appropriately as per supplier recommendations.
C	6	1.5.8 All standards used are within their expiration date.
C	2, 3	1.5.9 All standards are prepared <u>either gravimetrically or</u> using appropriate positive displacement pipettes or syringes.
C	3	1.5.10 Working standards are made up from primary standard by dilution with the toxin-free, extraction solvent (i.e., 100% methanol).
		1.6 Collection and Transportation of Samples
O	5, 1	1.6.1 Shellstock are collected in clean, waterproof, puncture resistant containers.
K	5, 1	1.6.2 Samples are appropriately labeled with the collector's name, type of shellstock, the harvest area, and time and date of collection.
C	5, 1	1.6.3 Immediately after collection, shellstock samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory.

K	2	1.6.4 Time from collection to initiation of the extraction should not exceed 24 hours. However, if significant delays are anticipated or if they occur, the laboratory has an appropriate contingency plan in place to handle the samples. For samples shipped live in accordance with 1.6.3, the contingency plan ensures samples remain within allowable temperature tolerances and animals are alive upon receipt. The contingency plan also addresses field and/or laboratory processing that ensures the integrity of the sample or extract until initiation of the assay. For example, samples are washed, shucked, drained and processed as follows: <ul style="list-style-type: none"> a. refrigerated or frozen until extracted; b. homogenized and frozen until extracted; or c. extracted, the supernatant decanted, and refrigerated or frozen until assayed.
C	2	1.6.5 Frozen shucked product or homogenates are allowed to thaw completely and all liquid is included as part of the sample before being processed further.
PART II – EXAMINATION OF SHELLFISH FOR DSP TOXINS		
		2.1 Preparation of Sample
C	2	2.1.1 At least 12 animals are used per sample or the laboratory has an appropriate contingency plan for dealing with non-typical species of shellfish (e.g., three (3) geoduck gut balls).
O	5	2.1.2 The outside of the shell is thoroughly cleaned with fresh water.
O	5	2.1.3 Shellstock are opened by cutting the adductor muscles.
O	5	2.1.4 The inside surfaces of the shells are rinsed with fresh water to remove sand and other foreign materials.
O	5	2.1.5 Shellfish meats are removed from the shell by separating the adductor muscles and tissue connecting at the hinge.
C	5	2.1.6 Damage to the body of the mollusk is minimized in the process of opening.
O	5	2.1.7 Shucked shellfish are drained on a #10 mesh sieve or equivalent without layering for five (5) minutes.
K	5	2.1.8 Pieces of shell and drainage are discarded.
C	2, 5	2.1.9 Drained meats or previously cooled/refrigerated shucked meats and their drip loss liquid or thawed homogenates with their freeze-thaw liquid are blended at high speed until homogenous (60-120 seconds).

		2.2 Sample Extraction
K	2	2.2.1 Sample homogenates are extracted as soon as possible (preferably the same day) or stored in the freezer at -10 °C or below.
C	3	2.2.2 Two (2.00) ± 0.05 g of homogenized sample is weighed into a 50 mL polypropylene centrifuge tube and subsequently extracted.
C	3	2.2.3 The sample homogenate is extracted with 9 mL of 100% methanol and vortexed to mix.
K	3	2.2.4 The sample homogenate/extract mixture is centrifuged for 10 minutes at 2000 x g and the supernatant decanted into a clean <u>container (e.g. new polypropylene tube or glass vial)</u> .
C	3	2.2.5 The tissue pellet is reextracted with nine (9) mL of 100% methanol and homogenized to mix.
K	3	2.2.6 The sample homogenate/extract mixture is centrifuged for 10 minutes at 2000 x g and the supernatant combined with the supernatant in 2.2.4.
K	3	2.2.7 The total extract volume <u>in the polypropylene tube</u> is adjusted to 20 mL with 100% methanol.
K	3	2.2.8 The crude extract is hydrolyzed or stored in the freezer at < -20 <u>10</u> °C.
		2.3 Sample Hydrolysis and Cleanup
K	3	2.3.1 A two (2) mL aliquot of the sample extract is transferred to an <u>appropriately sized* 16 × 100 mm</u> glass tube with a phenolic PTFE lined screw cap using a positive displacement pipette or syringe. <u>*Note: A 16 x 100 mm tube will have sufficient volume to perform the hydrolysis and hexane wash steps and fit in a standard 15 mL centrifuge tube adaptor.</u>
K	3	2.3.2 The sample extract is hydrolyzed by adding 250 µL of 2.5 M NaOH and the sample is homogenized with a vortex mixer for 30 seconds.
C	3	2.3.3 Sample tube caps are securely fastened to prevent extract loss, and the weight of the sample tube is recorded. The sample tube is heated at 76 °C for 40 minutes, then allowed to cool to room temperature, dried, and re-weighed. If the weight has dropped by more than 0.1 g, lost volume is replaced using 100% MeOH.
K	3	2.3.4 Samples are neutralized with 250 µL of 2.5 M HCL and vortexed to mix.
K <u>O</u>	3	2.3.5 Nonpolar lipids are removed by adding seven (7) mL of hexanes to the hydrolyzed sample extract and vortexing to mix (2.3.5 – 2.3.7 <u>OptionalPreferred</u>).
K	3	2.3.6 The sample extract/hexane mixture is partitioned by centrifuging for 10 minutes at 2000 x g (2.3.5 – 2.3.7 <u>OptionalPreferred</u>).

K	3	2.3.7 The hexane layer is removed with a glass pipette and one (1) mL of the hydrolyzed methanolic extract is removed and filtered into an LC-MS certified -glass <u>autosampler</u> vial using a 0.2 µm PTFE syringe tip filter (2.3.5 – 2.3.7 Optional <u>Preferred</u>).																														
K	<u>2.3</u>	2.3.8 The cleaned-up extract is loaded into the autosampler immediately for analysis.																														
2.4 Analysis																																
C	3	<p>2.4.1 Analytes are detected in standards and samples using the mass transitions in the table (negative ion mode).</p> <table border="1" style="margin-left: auto; margin-right: auto;"> <thead> <tr> <th>Compound</th> <th>Q1 (m/z)</th> <th>Q3 (m/z)</th> </tr> </thead> <tbody> <tr> <td>OA</td> <td>-803.5</td> <td>-255.2</td> </tr> <tr> <td>OA</td> <td>-803.5</td> <td>-113.1</td> </tr> <tr> <td>OA</td> <td>-803.5</td> <td>-151.1</td> </tr> <tr> <td>DTX2</td> <td>-803.5</td> <td>-255.2</td> </tr> <tr> <td>DTX2</td> <td>-803.5</td> <td>-113.1</td> </tr> <tr> <td>DTX2</td> <td>-803.5</td> <td>-151.1</td> </tr> <tr> <td>DTX1</td> <td>-817.5</td> <td>-255.2</td> </tr> <tr> <td>DTX1</td> <td>-817.5</td> <td>-113.1</td> </tr> <tr> <td>DTX1</td> <td>-817.5</td> <td>-151.1</td> </tr> </tbody> </table>	Compound	Q1 (m/z)	Q3 (m/z)	OA	-803.5	-255.2	OA	-803.5	-113.1	OA	-803.5	-151.1	DTX2	-803.5	-255.2	DTX2	-803.5	-113.1	DTX2	-803.5	-151.1	DTX1	-817.5	-255.2	DTX1	-817.5	-113.1	DTX1	-817.5	-151.1
Compound	Q1 (m/z)	Q3 (m/z)																														
OA	-803.5	-255.2																														
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DTX1	-817.5	-113.1																														
DTX1	-817.5	-151.1																														
K	3	2.4.2 Other system parameters such as collision energy are optimized for the specific system using standards before analysis.																														
C	3	2.4.3 A standard calibration curve of at least six (6) concentrations is performed before and after each set of samples <u>at the beginning of each run. An additional curve is required if a run lasts longer than 24 hours.</u>																														
K	3	2.4.4 Five (5) µL of extract is injected for analysis.																														
K	<u>2.12</u>	2.4.5 Samples are stored in the sample compartment of the autosampler at ≤ 10°C during analysis. Otherwise the samples must be analyzed within 24 hours if the autosampler is held at room temperature.																														
K	3	2.4.6 A column heater is used and the temperature is maintained at 40 °C, <u>with a tolerance as specified by the manufacturer</u> , during the analysis.																														
C	3	2.4.7 An Acquity UPLC BEH C18 1.0 × 150 mm, 1.7 µm particle size (or equivalent) analytical column is used for analyte separation																														
C	3	2.4.8 Analytes are separated on the LC column using gradient elution.																														

K	2	2.4.9 The column is stored following the manufacturer’s instructions when not in use.
K	2	2.4.10 Dead volume in the system is minimized by the use of short lengths of connecting tubing of small internal diameter between the sample injector and the column and between the column and detector.
C	3	2.4.11 Procedural <u>A matrix or procedural b</u> lanks (i.e. NRC CRM Zero-Mus or equivalent negative matrix, or methanol carried through sample preparation process at the same time as the samples) should be analyzed before and after extracted samples <u>included in each analytical batch.</u>

2.5 System Suitability		
C	3	2.5.1 Each calibration curve should be derived from at least six (6) calibration points and the linear regression of the combined curves should yield a correlation coefficient (R^2) ≥ 0.98 . Results are recorded and records are maintained.
C	3	2.5.2 If a calibration curve yields a correlation coefficient ≤ 0.98 , or if non-linearity is visually observed, or if the variation in the slopes between the first and second calibration curves exceeds 25%, a new calibration curve is prepared and samples are reanalyzed.
C	3	2.5.3.1 The retention time of analytes in all matrix solution should be within 3% that of the toxin standards. <u>all samples are within 3% that of one of the intermediate toxin standards, measured from the apex of the peak.</u>
C	3	2.5.4.2 Chromatographic separation must be sufficient for resolving OA and DTX2. Peak resolution (R_s) of OA/DTX2 should ≥ 1 when calculated using the equation below (RT is retention time and W is peak width <u>at baseline; Peak 1 is OA and Peak 2 is DTX2</u>). $R_s = 2 \times (RT_2 - RT_1) / (W_1 + W_2)$
K	2,3	2.5.5.3 Each chromatographic peak must be defined by at least 10 data points.
C	3	<u>2.5.4 Reagent blanks (methanol) are analyzed after the high calibration standard, periodically (as determined by the laboratory's internal verification), and after fortified samples to ensure that analyte carryover is not occurring. Analyte carryover is defined as a confirmed peak > LOD.</u>
C	3	<u>2.5.5 To confirm the presence of each DST, two (2) mass transitions must be observed above the limit of detection (LOD). The transition yielding the highest signal-to-noise ratio (S/N) is used for quantitation (i.e. 817.5 → 255.2 for DTX-1, 803.5 → 255.2 for OA and DTX-2). The transition yielding the second highest S/N is used for confirmation. The S/N of the peak used for confirmation is ≥ 3.</u>
C	3	<u>2.5.6 The ratio of the abundance of the confirmation ion transition to the quantitation ion transition is calculated for each toxin. These ion ratios must be within $\pm 20\%$ of that of the toxin standards in order to confirm toxin identity.</u>
C	3	<u>2.5.7 When ≥ 10 samples are analyzed, analysts must show that the calibration has not significantly drifted using an option provided below:</u>

		<u>OPTION 1 – USE OF BRACKETING CALIBRATION CURVES</u>
<u>C</u>	<u>3</u>	<u>2.5.7.1 A second standard curve is analyzed at the end of the analytical sequence and the averaged peak areas are used for the linear regression.</u>
<u>C</u>	<u>3</u>	<u>2.5.7.2 The linear regression of the averaged calibration curves must yield an R² ≥ 0.98. Results are recorded and records are maintained.</u>
<u>C</u>	<u>3</u>	<u>2.5.7.3 A new calibration curve is prepared, and samples are reanalyzed, if any of the following are observed:</u> <u>a) The average of bracketing calibration curves yields an R² < 0.98.</u> <u>b) The difference in the slope between bracketing calibration curves exceeds 25%.</u> <u>c) The difference in retention times of the standards in the bracketing standard curves exceeds 3%.</u>
		<u>OPTION 2 – USE OF A CONTINUING CALIBRATION VERIFICATION (CCV) STANDARD</u>
<u>C</u>	<u>3</u>	<u>2.5.7.4 The linear regression of the single calibration curve must yield an R² ≥ 0.99.</u>
<u>C</u>	<u>3</u>	<u>2.5.7.5 A continuing calibration verification (CCV) standard, matching on of the intermediate standards from the calibration curve, is analyzed after every 10 samples and at the end of a run.</u>
<u>C</u>	<u>3</u>	<u>2.5.7.6 A new calibration curve is prepared, and samples are reanalyzed, if any of the following are observed:</u> <u>a) The calibration curve yields an R² < 0.99.</u> <u>b) The retention time of a CCV sample exceeds 3% of the corresponding standard.*</u> <u>c) The peak area of a CCV sample exceeds ± 15% compared to the corresponding standard in the calibration curve.*</u> <u>* Samples immediately preceding and post the failed CCV shall be reanalyzed with a new standard curve.</u>
<u>C</u>	<u>3</u>	<u>2.5.8 Repeated injections of calibration or control samples at a concentration near the action level agree within ± 10% (as determined through the use of the coefficient of variation).</u>
<u>€</u>	<u>3</u>	<u>2.5.6 A new calibration curve is performed, or one mid-point calibration standard is analyzed, at least every 10 samples to ensure that no retention time shifts or loss in signal intensity has occurred.</u>
<u>K</u>	<u>2</u>	<u>2.5.7 Peak asymmetry must be <0.9 or >1.3.</u>

€	3	2.5.8 Reagent blanks (methanol) are analyzed after the high calibration standard and periodically after fortified samples to insure that analyte carryover is not occurring.
€	2	2.5.9 Repeated injections of calibrated standards/samples agree within ± 5% (as determined through the use of the coefficient of variation).
€	3	2.5.10 To confirm the presence of each DST, two (2) mass transitions must be observed above the limit of detection (LOD). The transition yielding the highest signal to noise ratio (S/N) is used for quantitation (i.e., 817.5 → 151.1 for DTX-1, 803.5 → 151.1 for OA and DTX-2). The transition yielding the second highest S/N is used for confirmation. The S/N of the peak used for confirmation is ≥ 3.
€	3	2.5.11 The ratio of the abundance of the quantitative ion transition to the confirmation ion transition is calculated for each toxin. These ion ratios must be within ± 20% of that of the toxin standards in order to confirm toxin identity.
		2.6 Calculation of Sample Toxicity

C	4, 15, 3, 11	<p>2.6.1 The toxicity of the individual toxins is calculated as follows:</p> $\frac{\mu\text{g}}{\text{g}} \text{ toxin} = C \times \frac{V}{W} \times \text{Hyd} \times \text{ReTx}$ <p>where:</p> <p>C = the concentration in µg/ml of the extract injected, determined using the standard curve</p> <p>V = total volume of homogenate and extraction solvent mL)</p> <p>W = weight (g) of tissue homogenate extracted</p> <p>Hyd = dilution factor for hydrolysis (1.25) Hyd = dilution factor for hydrolysis (1.25)</p> <p>ReTx = relative toxicity of toxin vs. Okadaic Acid ReTx = relative toxicity of toxin vs. Okadaic Acid</p> <p><u>Relative Toxicity Values</u> Relative Toxicity Values</p> <table border="1"> <thead> <tr> <th>Toxin</th> <th>ReTx</th> </tr> </thead> <tbody> <tr> <td>OA</td> <td>1</td> </tr> <tr> <td>DTX1</td> <td>1</td> </tr> <tr> <td>DTX2</td> <td>0.60.6</td> </tr> </tbody> </table> <p>The individual toxicities for each toxin are summed to obtain the overall sample toxicity in ppm OA equivalents. The individual toxicities for each toxin are summed to obtain the overall sample toxicity in ppm OA equivalents</p>	Toxin	ReTx	OA	1	DTX1	1	DTX2	0.60.6
Toxin	ReTx									
OA	1									
DTX1	1									
DTX2	0.60.6									
C	129	<p>2.6.2 Any value at or above 0.0 0.16 ppm OA equivalents OA equivalents (mg/kg or µg/g) of the sum of any analytes present is actionable. Shellfish Program Management is made aware of positive result. Laboratory action to identify positive result is _____ .</p>								
REFERENCES										
<p>1. American Public Health Association. 1984. <i>Compendium for the Microbiological Examination of foods</i>, 2nd Edition. APHA, Washington D.C.</p>										
<p>2. Good Laboratory Practice. 21 CFR 58.</p>										
<p>3. Interstate Shellfish Sanitation Conference (ISSC), Proposal 17-103 Liquid Chromatography Tandem Mass Spectrometry (LC-MS/MS) Method for the Determination of Diarrhetic Shellfish Poisoning (DSP) Toxins in Shellfish</p>										
<p>4. Association of Official Analytical Chemists (AOAC). 1991. <i>Quality Assurance Principles for Analytical Laboratories</i>. AOAC, Arlington, VA.</p>										
<p>5. American Public Health Association. 1970. <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i>, 4th Edition. APHA, Washington, D.C.</p>										

6. Consult reference standard product literature.
7. APHA/WEF/AWWA. 1992. <i>Standard Methods for the Examination of Water and Wastewater</i> , 18 th Edition. APHA, Washington, D.C.
8. American Public Health Association. 1992. <i>Standard Methods for the Examination of Dairy Products</i> , 16 th Edition. APHA, Washington, D.C.
9. U.S. Food and Drug Administration (FDA) and Interstate Shellfish Sanitation Conference (ISSC). 2015. <i>NSSP Guide for the Control of Molluscan Shellfish</i> . FDA/ISSC, Washington, D.C. and Columbia, S.C.
10. U.S. Department of Commerce. 1976. NBS Monograph 150. U.S. Department of Commerce, Washington, D.C.
<u>11. The EFSA Journal. 2009. Scientific Opinion of the Panel on Contaminants in the Food Chain on a request from the European Commission on Marine Biotoxins in Shellfish – Summary on regulated marine biotoxins. 1306, 1-23.</u>
<u>12. Deeds, J.R. and M.D. Celiz, Personal Communication, Addendum to proposal 19-136 DSP LC MS/MS Additional Ruggedness Testing; effect of the sample storage temperature during analysis (24 hrs), Email Received by ISSC Checklist Subcommittee 2/18/2021.</u>

LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
DIARRHETIC SHELLFISH POISON (DSP) COMPONENT: PARTS I AND II	
A. Results Total # of Critical (C) Nonconformities _____ Total # of Key (K) Nonconformities _____ Total # of Critical, Key, and Other (O) Nonconformities _____	_____ _____ _____
B. Criteria for Determining Laboratory Status of the DSP Component <ol style="list-style-type: none"> 1. Conforms Status: The DSP component of this Laboratory is in conformity with NSSP requirements if all of the following apply. <ol style="list-style-type: none"> a. No Critical nonconformities. b. and <6 Key nonconformities. c. and <12 Total nonconformities. 2. Provisionally Conforms Status: The DSP component of this laboratory is determined to be provisionally conforming to NSSP requirements if all of the following apply. <ol style="list-style-type: none"> a. the number of critical nonconformities is ≥ 1 but < 4. b. and < 6 Key nonconformities. c. and < 12 Total nonconformities. 3. Does Not Conform Status: The DSP component of this laboratory is not in conformity with NSSP requirements when any of the following apply. <ol style="list-style-type: none"> a. The total # of Critical nonconformities is ≥ 4. b. or the total # of Key nonconformities is ≥ 6. c. or the total # of Critical, Key, or Other is ≥ 12. 	
C. Laboratory Status (circle appropriate) <div style="text-align: center;"> Does Not Conform – Provisionally Conforms – Conforms </div>	
Acknowledgement by Laboratory Director/Supervisor: All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____. Laboratory Signature: _____ Date: _____ LEO Signature: _____ Date: _____	

Addendum to Proposal 19-136:

Single Laboratory Validation (SLV) Protocol for Submission to the Interstate Shellfish Sanitation Conference (ISSC) For Method Approval

Submitter: Jonathan Deeds, Ph.D.

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jonathan.deeds@fda.hhs.gov

Additional Ruggedness Testing: Effect of the sample storage temperature during analysis (24 hrs)

To assess the effect of sample storage temperature during analysis on accuracy/trueness, two sub-samples from each of 10 extracts from previously spiked samples, representing two different matrix sources (5 samples each from matrix sources A and B), each spiked at 5 different concentrations bracketing the regulatory guidance level (8, 12, 16, 24, and 32 $\mu\text{g}/100\text{ g}$), were tested at 10°C and also at room temperature (approx. 24°C) after storage at room temperature for 24 hrs. After hydrolysis and hexane washing, each sample was filtered, as described previously, into two LC-vials. One set was analyzed using a refrigerated sample injector set to 10°C, while the second set was stored at room temperature (approx. 24 °), protected from light, for 24 hrs, then injected using the same sample injector with the refrigeration turned off. This entire procedure was repeated on separate days so that in total 20 samples were tested at 10°C and at 24°C after storage for 24 hrs. The data handling procedures outlined in the Marine Biotxin and Non-MPN Based Microbiological Methods SOP for Ruggedness were used for reporting these results. Data for the ruggedness testing of the sample storage temperature during analysis are presented in tables 25, 26, and 27. The variance ratio showed no significant differences between the two treatments for any of the three toxins, and no significant differences were found in the measured concentrations if injected at 10°C or at room temperature after storage for 24 hrs. We conclude from this data that during analysis, finished samples can be stored at room temperature for up to 24 hrs. without significantly affecting the analytical results.

Table 25. Ruggedness Testing for Sample Storage Temperature During Analysis for DTX1 in Clam

DTX1		10°C	Room Temp (ca. 24°C)
Sample	Spiked Concentration (µg/100g)	Determined Concentration (µg/100g)	Determined Concentration (µg/100g)
1	8	6.984	7.413
2	12	10.835	10.784
3	16	14.455	14.093
4	24	21.329	20.961
5	32	27.770	28.106
6	8	7.144	6.889
7	12	11.219	10.575
8	16	14.870	14.364
9	24	22.573	21.815
10	32	30.958	30.344
11	8	7.215	7.073
12	12	10.571	10.225
13	16	14.086	13.856
14	24	20.865	22.201
15	32	27.650	28.030
16	8	6.933	6.834
17	12	10.295	10.516
18	16	14.266	14.350
19	24	21.179	21.666
20	32	29.179	38.408
Skewness			
		0.47	0.82
Variance			
		64.7	82.4
Variance Ratio =1.3, no significant difference			
Paired t-test (two-tailed): P=0.405, no significant difference			
Mean of differences: 0.84; 95% confidence interval: -0.09-1.8			

Table 26. Ruggedness Testing for Sample Storage Temperature During Analysis for DTX2 in Clam

DTX2		10°	Room Temp (ca. 24°C)
Sample	Spiked Concentration (µg/100g)	Calculated Concentration (µg/100g)	Calculated Concentration (µg/100g)
1	8	7.215	7.006
2	12	10.268	10.253
3	16	13.664	13.326
4	24	20.378	19.954
5	32	27.649	26.761
6	8	7.236	6.590
7	12	10.281	9.823
8	16	13.925	13.553
9	24	21.863	20.364
10	32	30.780	28.513
11	8	6.430	6.489
12	12	9.428	9.308
13	16	12.624	12.949
14	24	18.985	20.738
15	32	24.391	26.468
16	8	6.395	6.396
17	12	9.325	9.644
18	16	12.226	13.653
19	24	19.168	20.313
20	32	26.528	34.255
Skewness		0.592	0.714
Variance		59.4	69.8
Variance Ratio =1.2, no significant difference			
Paired t-test (two-tailed): P=0.410, no significant difference			
Mean of differences: 1.1; 95% confidence interval: 0.30-1.9			

Table 27. Ruggedness Testing for Sample Storage Temperature During Analysis for OA in Clam

OA		10°	Room Temp (ca. 24°C)
Sample	Spiked Concentration (µg/100g)	Calculated Concentration (µg/100g)	Calculated Concentration (µg/100g)
1	8	7.236	7.018
2	12	10.856	10.399
3	16	14.356	13.736
4	24	22.198	20.748
5	32	29.823	27.436
6	8	6.954	6.853
7	12	11.179	10.421
8	16	15.896	14.149
9	24	23.409	21.384
10	32	33.059	29.430
11	8	6.578	6.945
12	12	9.645	10.685
13	16	13.321	14.136
14	24	20.296	22.180
15	32	25.723	27.905
16	8	6.521	6.621
17	12	9.813	10.376
18	16	13.580	14.253
19	24	20.433	21.889
20	32	28.225	37.651
Skewness			
		0.542	0.784
Variance			
		70.2	79.4
Variance Ratio =1.1, no significant difference/variance homogeneous			
Paired t-test (two-tailed): P=0.667, no significant difference			
Mean of differences: 1.6; 95% confidence interval: 0.63-2.6			

PUBLIC HEALTH SERVICE U.S. FOOD AND DRUG ADMINISTRATION OFFICE OF FOOD SAFETY SHELLFISH AND AQUACULTURE POLICY BRANCH 5100 PAINT BRANCH PARKWAY COLLEGE PARK, MD 20740-3835 TEL. 240- 402-2151/2055/4960 FAX 301-436-2601		
SHELLFISH LABORATORY EVALUATION CHECKLIST		
LABORATORY:		
ADDRESS:		
TELEPHONE:		FAX:
EMAIL:		
DATE OF EVALUATION:	DATE OF REPORT:	LAST EVALUATION:
LABORATORY REPRESENTED BY:		TITLE:
LABORATORY EVALUATION OFFICER:		SHELLFISH SPECIALIST:
		REGION:
OTHER OFFICIALS PRESENT:		TITLE:
Items which do not conform are noted by:		Conformity it noted by a “√”
C- Critical K - Key O - Other NA- Not Applicable		
Check the applicable analytical methods:		
<input type="checkbox"/> Multiple Tube Fermentation Technique for Seawater (APHA)[PART II]		
<input type="checkbox"/> Multiple Tube Fermentation Technique for Seawater using MA-1 [PART II]		
<input type="checkbox"/> Membrane Filtration Technique for Seawater using mTEC [PART II]		
<input type="checkbox"/> Multiple Tube Fermentation Technique for Shellfish Meats (APHA)[PART III]		
<input type="checkbox"/> Standard Plate Count for Shellfish Meats [PART III]		
<input type="checkbox"/> Elevated Temperature Coliform Plate Method for Shellfish Meats [PART III]		
<input type="checkbox"/> Male Specific Coliphage for Soft-shelled Clams and American Oysters [PART III]		

PART 1 - QUALITY ASSURANCE

CODE	REF.	ITEM
K	8, 11	1.1 Quality Assurance (QA) Plan
		1.1.1 Written Plan (Check those items which apply.)
		a. Organization of the laboratory.
		b. Staff training requirements.
		c. Standard operating procedures.
		d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		e. Laboratory safety.
		f. Internal performance assessment.
		g. External performance assessment.
C	8	1.1.2 QA Plan Implemented.
K	11	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s) _____
		1.2 Educational/Experience Requirements
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three months of experience in laboratory sciences.
		1.3 Work Area
O	8,11	1.3.1 Adequate for workload and storage.
K	11	1.3.2 Clean, well-lighted.
K	11	1.3.3 Adequate temperature control.
O	11	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
		1.4 Laboratory Equipment
O	9	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	1.4.4 pH meter is calibrated daily or with each use Results are recorded and records maintained.
K	11	1.4.5 <u>The pH meter manufacturer instructions are followed for calibration, or A</u> minimum of two standard buffer solutions is used to calibrate the pH meter. <u>If the calibration sequence of standard buffer solutions is not stipulated by the manufacturer, T</u> the first must be near the electrode isopotential point (pH 7) <u>and T</u> the second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.

O	8,15	1.4.6	Electrode acceptability is determined daily or with each use by the millivolt procedure or through determination of the slope. <i>(Circle the method used.)</i>
K	9	1.4.7	Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13	1.4.8	Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11	1.4.9	Refrigerator temperature(s) are monitored at least once daily on workdays. Results are recorded and records maintained.
K	1	1.4.10	Refrigerator temperature is maintained at 0 to 4°C.
C	9	1.4.11	The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11	1.4.12	Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9	1.4.13	Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11	1.4.14	Temperature of the water bath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9	1.4.15	The thermometers used in the water_bath are graduated in at least 0.1°C increments.
C	13	1.4.16	The water_bath has adequate capacity for workload.
K	9	1.4.17	The level of water in the water bath covers the level of liquid in the incubating tubes.
K	8, 11	1.4.18	Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4	1.4.19	All working thermometers are appropriately immersed.
C	29-	1.4.20	Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	11	1.4.21	A mercury-in-glass standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9	1.4.22	Standards thermometers are checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination_____.
C	29	1.4.23	Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of ≤ ±0.05°C are used as the laboratory standards thermometer. <i>(Circle the thermometer type used.)</i>
K	13	1.4.24	Incubator and water bath working thermometers are checked annually against the standards thermometer at the temperatures at which they are used. Results are recorded and records maintained.
O	11	1.4.25	Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing			
O	9	1.5.1	Utensils and containers are clean borosilicate glass, stainless steel, or other noncorroding materials.
K	9	1.5.2	Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.

K	9	1.5.3	Sample containers are made of glass or some other inert material.
O	9	1.5.4	Dilution bottles and tubes are made of borosilicate glass or plastic and closed with rubber stoppers, caps or screw caps with nontoxic liners.
K	9	1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9	1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1 mL used to deliver 0.1 mL aliquots.
K	9	1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9	1.5.8	In washing reusable pipettes, a succession of at least three fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2	1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11	1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination			
K	9	1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8	1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	11, 30	1.6.3	The autoclave provides a sterilizing temperature of 121 ± 2°C as determined for each load using a calibrated maximum registering thermometer. As an alternative, an appropriate temperature monitoring device is used in place of the maximum registering thermometer when these are unavailable due to the ban on mercury.
K	11	1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16	1.6.5	The autoclave standards thermometer is checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory or is checked in-house at the steam point (100°C) if it has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination _____
K	1	1.6.6	Working autoclave thermometers are checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11	1.6.7	Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11	1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13	1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11	1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of ≥ 160 to 180 °C.

K	9		1.6.11 A thermometer capable of determining temperatures accurately in the range of ≥ 160 to 180°C <u>accurately</u> is used to monitor the operation of the hot-air sterilizing oven.
K	13		1.6.12 Records of temperatures and exposure times are maintained for the operation of the hot-air sterilizing oven during use.
K	11		1.6.13 Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Records are maintained.
K	11		1.6.14 Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C .
C	1		1.6.15 The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1		1.6.16 The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9		1.6.17 Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9		1.6.18 Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2 hours.
C	2		1.6.19 The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2		1.6.20 The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18		1.6.21 Hardwood applicator transfer sticks are properly sterilized. Method of sterilization _____
C	2		1.6.22 The sterility of the hardwood applicator transfer sticks is checked routinely. Results are recorded and the records maintained.
O	13		1.6.23 Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation			
K	3, 5		1.7.1 Media is commercially dehydrated except in the case of medium A-1 which must be prepared from the individual components and modified MacConkey Agar which may be prepared from its components.
K	11		1.7.2 Media is prepared according to manufacturer's instructions.
O	11		1.7.3 Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11		1.7.4 Dehydrated media are labeled with date of receipt and date opened.
C	12		1.7.5 Caked or expired media or media components are discarded.
C	11		1.7.6 Reagent water is distilled or deionized (circle appropriate choice), tested monthly and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) or is less than 2.0 $\mu\text{Siemens/cm}$ conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11		1.7.7 Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level ($< 0.1 \text{ mg/L}$). Results are recorded and the records maintained. Specify method of determination _____.
K	11		1.7.8 Reagent water contains $< 100 \text{ CFU/mL}$ as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11		1.7.9 Media prepared from commercial dehydrated components sterilized according to the manufacturer's instructions.
K	9		1.7.10 The volume and concentration of media in the tube are suitable for the amount of sample inoculated.

C	11		1.7.11	Total time of exposure of sugar broths to autoclave temperatures does not exceed 45 minutes.
C	1		1.7.12	Media sterility is determined for each load sterilized. Results are recorded and the records maintained.
C	1		1.7.13	Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is made from its individual components.
O	9		1.7.14	Sterile phosphate buffered dilution water is used as the sample diluent.
K	11		1.7.15	The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media				
K	9		1.8.1	Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11		1.8.2	Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13		1.8.3	Stored media are labeled with the storage expiration date or the sterilization date.
K	9		1.8.4	Storage of prepared culture media at room temperature does not exceed 7 days.
K	2		1.8.5	Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed 1 month.
K	11		1.8.6	Storage under refrigeration of prepared culture media with screw-cap closures does not exceed 3 months.
K	17		1.8.7	All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES				
2.1 Collection and Transportation of Samples				
C	11		2.1.1	Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1		2.1.2	Samples are identified with collectors name, harvest area, sampling station, time and date of collection.
C	9		2.1.3	Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1		2.1.4	A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9		2.1.5	Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.
2.2 Bacteriological Examination of Seawater by the APHA MPN				
C	9		2.2.1	Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
C	2		2.2.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	9		2.2.3	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9		2.2.4	In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).

C	6	2.2.5	In a single dilution series not less than 12 tubes are used (for depuration at least 5 tubes are used).
C	6	2.2.6	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
K	9	2.2.7	Inoculated tubes are incubated in air at 35 ± 0.5 °C.
C	2	2.2.8	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
K	9	2.2.9	Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
2.3 Confirmed Test for Seawater by APHA MPN			
C	9	2.3.1	Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9	2.3.2	EC medium is used as the confirmatory medium for fecal coliforms.
C	2	2.3.3	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11	2.3.4	Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. (Circle the method of transfer.)
C	9	2.3.5	BGB tubes are incubated at 35 ± 0.5 °C.
K	9	2.3.6	BGB tubes are read after 48 ± 3 hours of incubation.
C	9	2.3.7	EC tubes are incubated in a circulating water bath maintained at 44.5 ± 0.2 °C.
C	9	2.3.8	EC tubes are read after 24 ± 2 hours of incubation.
C	9	2.3.9	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.4 Computation of Results – APHA MPN			
K	9	2.4.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7	2.4.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.4.3	Results are reported as MPN/100 mL of sample.
2.5 Bacteriological Examination of Seawater by the MA-1 Method			
C	5	2.5.1	A-1 medium complete is used in the analysis.
C	2, 31	2.5.2	A-1 medium without salicin is used in the analysis. Comparability testing

			supports use of A-1 medium without salicin. Study records are available
C	5	2.5.3	A-1 medium sterilized for 10 minutes at 121°C.
C	2	2.5.4	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	9	2.5.5	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds) before inoculation.
C	9	2.5.6	In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	2.5.7	In a single dilution series at least 12 tubes are used.
C	6	2.5.8	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	2.5.9	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and water bath incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	2.5.10	Inoculated tubes are placed in an air incubator at 35 ± 0.5°C for 3 ± 0.5 hours of resuscitation.
C	5	2.5.11	After 3 ± 0.5 hours resuscitation at 35°C, inoculated tubes are incubated at 44.5 ± 0.2°C in a circulating water bath for the remainder of the 24 ± 2 hours.
C	5	2.5.12	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
		2.6 Computation of Results – APHA MPN	
K	9	2.6.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	2.6.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.6.3	Results are reported as MPN/100 mL of sample.
		2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment	
C	23, 24	2.7.1	When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at 44.5 ± 0.5°C under any loading capacity.
C	23	2.7.2	When using a water bath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	2.7.3	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	2.7.4	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	2.7.5	Colonies are counted with the aid of magnification.
C	11, 23	2.7.6	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of 0.45 µm and certified by the manufacturer for fecal coliform analyses.
C	2	2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.

C	2		2.7.8 When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison testing implemented. The results are recorded and this record is maintained.
K	2, 11		2.7.9 New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2		2.7.10 The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2		2.7.11 Membrane filters which are beyond their expiration date are not used.
O	11		2.7.12 Forceps tips are clean.
O	11		2.7.13 Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.
K	11		2.7.14 Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11		2.7.15 If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11		2.7.16 Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11		2.7.17 Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26		2.7.18 A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11		2.7.19 The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2		2.7.20 Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
2.8 Media Preparation and Storage – MF using mTEC Agar			
K	11		2.8.1 Phosphate buffered saline is used as the sample diluent and filter funnelrinse.
C	11		2.8.2 The phosphate buffered saline is properly sterilized.
K	23		2.8.3 A sufficient amount of medium (4-5 mL) is used in each plate.
O	11		2.8.4 Refrigerated prepared plates are stored for no more than 2 weeks in sealed plastic bags or containers to minimize evaporation.
2.9 Sample Analyses - MF using mTEC Agar			
C	24		2.9.1 mTEC agar is used.
C	2		2.9.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	23		2.9.3 The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration.
C	23		2.9.4 The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25		2.9.5 Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23		2.9.6 Sample volumes are filtered under vacuum.
K	26		2.9.7 The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26		2.9.8 The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.

C	23	2.9.9	The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11	2.9.10	Blanks are run at the beginning of filtration, after every 10 th aliquot, and at the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).
C	2, 11	2.9.11	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24	2.9.12	Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at 35 ± 0.5 °C for 2 hours of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5 °C for 24 ± 2 hours.
C	11, 23, 24	2.9.13	After <u>two</u> (2) hours of resuscitation at 35°C, the watertight, tightly sealed containers are transferred to a circulating water bath at 44.5 ± 0.2°C, submerged completely and incubated for 22-24 hours.
2.10 Computation of Results - MF using mTEC Agar			
C	23	2.10.1	All yellow, yellow-green, or yellow-brown colonies are counted.
C	23	2.10.2	Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as >80 x 100/the volume of sample filtered.
C	2, 11, 23	2.10.3	When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11	2.10.4	The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11	2.10.5	Results are reported as CFU/100 mL of sample.
PART III - SHELLFISH SAMPLES			
3.1 Collection and Transportation of Samples			
C	9	3.1.1	A representative sample of shellstock is collected.
K	9	3.1.2	Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9	3.1.3	Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9	3.1.4	Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1	3.1.5	Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
3.2 Preparation of Shellfish for Examination			
K	2,11	3.2.1	Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2	3.2.2	Blades of shucking knives are not corroded.
O	9	3.2.3	The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.

O	2		3.2.4	The faucet used for rinsing the shellstock does not contain an aerator.
K	9		3.2.5	Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9		3.2.6	Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	9		3.2.7	Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9		3.2.8	Shellstock are not shucked directly through the hinge.
C	9		3.2.9	Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9		3.2.10	At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.
K	9		3.2.11	A representative sample of at least 12 shellfish is used for the analysis.
K	2		3.2.12	The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
O	9		3.2.13	Sterile phosphate buffered dilution water is used as the sample diluent.
C	9		3.2.14	Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9		3.2.15	APHA <i>Recommended Procedures for the Examination of Sea Water And Shellfish</i> , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA				
C	9		3.3.1	Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. (Circle the medium used.)
C	2		3.3.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9		3.3.3	Immediately (within 2 minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9		3.3.4	No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
C	9		3.3.5	Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6		3.3.6	In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2		3.3.7	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9		3.3.8	Inoculated media are incubated at 35 ± 0.5°C.
K	10		3.3.9	Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA				
C	9		3.4.1	EC medium is used as the confirmatory medium.
C	2		3.4.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____

K	9, 11		3.4.3	Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. (<i>Circle the method of transfer.</i>)
C	9		3.4.4	EC tubes are incubated in a circulating water bath at 44.5 ± 0.2°C
K	9		3.4.5	EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9		3.4.6	The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.
3.5 Computation of Results for MPN Analyses				
K	9		3.5.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.
K	7		3.5.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9		3.5.3	Results are reported as MPN/100 grams of sample.
3.6 Standard Plate Count Method				
O	20		3.6.1	A standard plate count (SPC) analysis may be performed in conjunction with the analysis for fecal coliform organisms.
K	9		3.6.2	In the standard plate count procedure at least four plates are used, duplicates of two dilutions. One of the dilutions should produce colonies of 30 to 300 per plate.
K	2		3.6.3	Fifteen to 20 mL of tempered sterile plate count agar is used per plate.
C	9		3.6.4	Agar tempering bath maintains the agar at 44-46°C.
C	9		3.6.5	An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9		3.6.6	Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in 7 seconds) before plating.
C	9		3.6.7	Not more than <u>one (1)</u> mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11		3.6.8	Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9,21		3.6.9	Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four high.
K	9		3.6.10	Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1		3.6.11	A hand tally or its equivalent is used for accuracy in counting.
3.7 Computation of Results -SPC				
K	9		3.7.1	Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19		3.7.2	Colony counts are reported as CFU/g of sample.
3.8 Bacteriological Analysis of Shellfish Using the ETCP				
C	2,3		3.8.1	Prepared modified MacConkey agar is used on the day that it is made.
K	3		3.8.2	Double strength modified MacConkey agar is used.
C	3		3.8.3	Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3		3.8.4	Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50 °C until used.
K	2, 3		3.8.5	Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3		3.8.6	The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9		3.8.7	The sample homogenate is cultured within 2 minutes of blending.
C	2,3		3.8.8	Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.

K	3		3.8.9	Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2,3, 22		3.8.10	The container is gently swirled or slowly inverted once to mix the contents, which are subsequently distributed uniformly over six plates.
C	1		3.8.11	Media and diluent sterility are determined with each use. Results are recorded and the records maintained.
C	1		3.8.12	Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture _____ Negative control culture _____
C	3, 13		3.8.13	When solidified, the plates are placed inverted into an air incubator at 45.5 ± 0.5°C for 18 to 30 hours of incubation.
C	2		3.8.14	Plates are stacked no more than three high in the incubator.
C	2		3.8.15	Appropriately diluted pour plated process control cultures accompany each set of samples throughout incubation. The results are recorded and the records maintained. Positive process control _____ Negative process control _____
3.9 Computation of Results - ETCP				
K	11		3.9.1	Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1		3.9.2	A hand tally or its equivalent is used to aid in counting.
C	3, 6		3.9.3	All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3		3.9.4	Results are reported as CFU/100 grams of sample.
Bacteriological Examination of Soft-shelled Clams and American Oysters for Male Specific Coliphage (MSC)				
3.10 MSC Equipment and Supplies				
K	30		3.10.1	Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold 100 – 125 mL.
C	27, 28		3.10.2	The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	9		3.10.3	The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
C	27, 28		3.10.4	Sterile 0.22 µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1		3.10.5	The sterility of each lot of pre-sterilized syringes and syringe filters is determined. Results are recorded and records maintained.
K	1		3.10.6	The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28		3.10.7	The balance used provides a sensitivity of at least 10 mg (0.01g.).
C	27, 28		3.10.8	The temperature of the incubator used is maintained at 36 ± 1°C.
C	28		3.10.9	Sterile disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
3.11 MSC Media Preparation				
K	28		3.11.1	Media preparation and sterilization is according to the validated method.
<u>CK</u>	27, 28		3.11.2, 3.11.2	Bottom agar, double strength soft agar and growth broth are prepared from their individual components.
K	27, 28		3.11.3	Soft agar is prepared double strength in volumes of 2.5 mL.
C	27, 28		3.11.4	The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
O	27, 28		3.11.5	Storage of the bottom agar under refrigeration does not exceed 1 month.
K	27, 28		3.11.6	Unsterilized soft agar is stored at -20 °C -15C for up to 3 months.
K	27, 28		3.11.7	The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.

K	27, 28		3.11.8 Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	27, 28		3.11.9 Bottom agar plates are allowed to reach room temperature before use.
3.12 Preparation of the Soft-Shell Clams and American Oysters for MSC Analysis			
K	2,11		3.12.1 Shucking knives, scrub brushes, and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2		3.12.2 The blades of shucking knives are not corroded.
O	9		3.12.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2		3.12.4 The faucet used for rinsing the shellfish does not contain an aerator.
K	9		3.12.5 The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9		3.12.6 The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9		3.12.7 Immediately prior to shucking, the hands (or gloved hands) of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol.
C	9		3.12.8 Shellfish are not shucked through the hinge.
C	9		3.12.9 The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9		3.12.10 A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19		3.12.11 The sample is weighed to the nearest 0.1 gram.
3.143 MSC Sample Analysis			
C	28		3.13.1 E.coli Famp ATCC 700891 is the bacterial host strain used in this procedure.
K	27, 28		3.13.2 Host cell growth broth is tempered at $36 \pm 1^\circ\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	27, 28		3.13.3 Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^\circ\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	27, 28		3.13.4 After inoculation, the host cell growth broth culture is not shaken.
C	28		3.13.5 A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.
C	28		3.13.6 The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28		3.13.7 The elution mixture is homogenized at high speed for 180 seconds.
C	28		3.13.8 Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
C	28		3.13.9 The homogenized elution mixture is centrifuged for 15 minutes at $9000 \times g$ at 4°C.
C	27, 28		3.13.10 The supernatant is pipetted off, weighed, and the weight recorded.
C	27, 28		3.13.11 The supernatant is allowed to warm to room temperature about 20 to 30 minutes.
K	27, 28		3.13.12 The autoclaved soft agar is tempered and held at $51 \pm 1^\circ\text{C}$ throughout the period of sample analysis.
K	27, 28		3.13.13 Two hundred microliters (0.2 mL) of log phase host strain <i>E coli</i> is added to the tempering soft agar immediately prior to adding the sample supernatant.
K	27, 28		3.13.14 The sample supernatant is shaken or vortexed before being added to the tempering soft agar.
C	27, 28		3.13.15 2.5 mL of sample supernatant is added to each tube of tempering soft agar.
C	27, 28		3.13.16 The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28		3.13.17 The soft agar/sample supernatant/host cell mixture is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.

C	28		3.13.18 Ten (10) plates are used 2.5 mL per plate for a total of 25 mL of supernatant analyzed per sample.
K	27, 28		3.13.19 Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are and records maintained. Positive control _____
K	27, 28		3.13.20 Growth broth is used as the negative control or blank.
K	27, 28		3.13.21 Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2		3.13.22 A negative control plate is plated at the beginning and end of each set of samples analyzed.
K	27, 28		3.13.23 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28		3.13.24 All plates are incubated at 36 ± 1 °C for 18 ± 2 hours.
			3.154 Computation of Results -MSC
C	27		3.14.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 32		3.14.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten (10) plates, the count is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/ 100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as > 20,000 PFU/100 grams.
K	28		3.14.3 The formula used for determining the density of MSC in PFU/100 grams is: $(0.364) (N) (Ws)$, where N = total number of plaques counted on all 10 plates and Ws = weight of the supernatant used.
O	9		3.14.4 The MSC count is rounded off conventionally to give a whole number.

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LABORATORY STATUS		
LABORATORY	DATE	
LABORATORY REPRESENTATIVE:		
MICROBIOLOGICAL COMPONENT: (Part I-III)		
A. Results		
Total # of Critical (C) Nonconformities in Parts I-III	_____	
Total # of Key (K) Nonconformities in Parts I-III	_____	
Total # of Critical, Key and Other (O)	_____	
Nonconformities in Parts I-III		
B. Criteria for Determining Laboratory Status of the Microbiological Component:		
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <ul style="list-style-type: none"> a. The total # of Critical nonconformities is ≥ 4 or b. The total # of Key nonconformities is ≥ 13 or c. The total # of Critical, Key and Other is ≥ 18 <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but ≤ 3.</p>		
C. Laboratory Status (circle appropriate)		
Does Not Conform	Provisionally Conforms	Conforms
<p>Acknowledgment by Laboratory Director/Supervisor:</p> <p>All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before_____.</p> <p>Laboratory Signature: _____ Date:_____</p> <p>LEO Signature: _____ Date:_____</p>		

National Shellfish Sanitation Program (NSSP) Guide for the Control of Molluscan Shellfish: 2019 Revision

PUBLIC HEALTH SERVICE
U.S. FOOD AND DRUG ADMINISTRATION
OFFICE OF FOOD SAFETY
SHELLFISH AND AQUACULTURE POLICY BRANCH
5100 PAINT BRANCH PARKWAY 5001 CAMPUS DRIVE
COLLEGE PARK, MD 20740-3835
TEL. 240- 402-21514960/9258/2055/4960 FAX 301-436-2601

SHELLFISH LABORATORY EVALUATION CHECKLIST

LABORATORY:

ADDRESS:

TELEPHONE:

FAX:

EMAIL:

DATE OF EVALUATION:

DATE OF REPORT:

LAST EVALUATION:

LABORATORY REPRESENTED BY:

TITLE:

LABORATORY EVALUATION OFFICER:

SHELLFISH SPECIALIST:

REGION:

OTHER OFFICIALS PRESENT:

TITLE:

Items which do not conform are noted by:

Conformity it noted by a “√”

C- Critical K - Key O - Other NA- Not Applicable

Check the applicable analytical methods:

- | | |
|--------------------------|--|
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Seawater (APHA)[PART II] |
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Seawater using MA-1 [PART II] |
| <input type="checkbox"/> | Membrane Filtration Technique for Seawater using mTEC [PART II] |
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Shellfish Meats (APHA)[PART III] |
| <input type="checkbox"/> | Standard Plate Count for Shellfish Meats [PART III] |
| <input type="checkbox"/> | Elevated Temperature Coliform Plate Method for Shellfish Meats [PART III] |
| <input type="checkbox"/> | Male Specific Coliphage for Soft-shelled Clams and American Oysters <u>Shellfish Meats</u> [PART III] |
| <input type="checkbox"/> | Membrane Filtration Technique for Seawater using mTEC [Part II] |
| <input type="checkbox"/> | Membrane Filtration Technique for UV Treated Process Water using mEndo Agar LES [Part II] |
| <input type="checkbox"/> | Multiple Tube Fermentation Technique for Shellfish Meats (APHA) [Part III] |

National Shellfish Sanitation Program (NSSP) Guide for the Control of Molluscan Shellfish: 2019 Revision

PART 1 - QUALITY ASSURANCE		
CODE	REF.	ITEM
K	8, 11	1.1 Quality Assurance (QA) Plan
		1.1.1 Written Plan (Check those items which apply.)
		a. Organization of the laboratory.
		b. Staff training requirements.
		c. Standard operating procedures.
		d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		e. Laboratory safety.
		f. Internal performance assessment.
		g. External performance assessment.
C	8	1.1.2 QA Plan Implemented.
K	11	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s)
		1.2 Educational/Experience Requirements
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two (2) years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three (3) months of experience in laboratory sciences.
		1.3 Work Area
O	8,11	1.3.1 Adequate for workload and storage.
K	11	1.3.2 Clean, well-lighted.
K	11	1.3.3 Adequate temperature control.
O	11	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
		1.4 Laboratory Equipment
O	9	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	1.4.4 pH meter is calibrated daily or with each use Results are recorded and records maintained.
K	11	1.4.5 A minimum of two (2) standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	8,15	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt

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			procedure or through determination of the slope. (<i>Circle the method used.</i>)
K	9		1.4.7 Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13		1.4.8 Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11		1.4.9 Refrigerator temperature(s) are monitored at least once daily on workdays. Results are recorded and records maintained.
K	1		1.4.10 Refrigerator temperature is maintained at 0 to 4°C.
C	9		1.4.11 The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11		1.4.12 Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9		1.4.13 Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11		1.4.14 Temperature of the waterbath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9		1.4.15 The thermometers used in the waterbath are graduated in at least 0.1°C increments.
C	13		1.4.16 The waterbath has adequate capacity for workload.
K	9		1.4.17 The level of water in the waterbath covers the level of liquid in the incubating tubes.
K	8, 11		1.4.18 Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4		1.4.19 All working thermometers are appropriately immersed.
C	299		1.4.20 Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	11		1.4.21 A standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9		1.4.22 Standards thermometers are checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination _____.
C	299		1.4.23 Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of <u>≤at least</u> ±0.05°C are used as the laboratory standards thermometer. (<i>Circle the thermometer type used.</i>)
K	13		1.4.24 The accuracy of working thermometers is checked annually against the standards thermometer either at the temperatures at which they are used or by ice point determination. Results are recorded and records maintained.
O	11		1.4.25 Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing			
O	9		1.5.1 Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding materials.
K	9		1.5.2 Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.
K	9		1.5.3 Sample containers are made of glass or some other inert material.
O	9		1.5.4 Dilution bottles and tubes are made of borosilicate glass or plastic and closed

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			with rubber stoppers, caps, or screw caps with nontoxic liners.
K	9	1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9	1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1mL used to deliver 0.1 mL aliquots.
K	9	1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9	1.5.8	In washing reusable pipettes, a succession of at least three (3) fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2	1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11	1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination			
K	9	1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8	1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	3029, 3332, 3433	1.6.3	The autoclave provides sterilization conditions suitable to the load contents. Sterilization temperature range may be 119°C - 124°C as determined by the lab's equipment Quality Assurance Verification Testing and recommended practices from the media manufacturer. Sterilization is determined for each load using a working -maximum registering thermometer or an appropriate working temperature monitoring device.
K	11	1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16	1.6.5	The autoclave standards thermometer is checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory; or, is checked in-house at the steam point (100°C) if it has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination _____
K	1	1.6.6	Working autoclave thermometers are checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11	1.6.7	Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11	1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13	1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11	1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180°C.
K	9	1.6.11	A thermometer capable of determining temperatures accurately in the range of 160 to 180°C is used to monitor the operation of the hot-air sterilizing oven
K	13	1.6.12	Records of temperatures and exposure times are maintained for the operation of

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			the hot-air sterilizing oven during use.
K	11	1.6.13	Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Records are maintained.
K	11	1.6.14	Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C.
C	1	1.6.15	The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1	1.6.16	The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9	1.6.17	Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9	1.6.18	Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for <u>two (2)</u> hours.
C	2	1.6.19	The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2	1.6.20	The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18	1.6.21	Hardwood applicator transfer sticks are properly sterilized. Method of sterilization _____
C	2	1.6.22	The sterility of the hardwood applicator transfer sticks is checked routinely. Results are recorded and the records maintained.
O	13	1.6.23	Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation			
K	3, 5	1.7.1	Media is commercially dehydrated except in the case of medium -A-1 <u>medium</u> , which must be prepared from the individual components and modified MacConkey agar which may be prepared from its components.
K	11	1.7.2	Media is prepared according to manufacturer's instructions.
O	11	1.7.3	Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11	1.7.4	Dehydrated media are labeled with date of receipt and date opened.
C	12	1.7.5	Caked or expired media or media components are discarded.
C	11	1.7.6	Reagent water is distilled or deionized (circle appropriate choice), tested monthly, and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) or is less than 2.0 µSiemens/cm conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11	1.7.7	Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level (< 0.1 mg/L). Results are recorded and the records maintained. Specify method of determination _____.
K	11	1.7.8	Reagent water contains <100 CFU/mL as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11	1.7.9	Media prepared from commercial ly dehydrated components are sterilized according to the manufacturer's instructions.
K	9	1.7.10	The volume and concentration of media in the tube are suitable for the amount of sample inoculated.
C	11	1.7.11	Total time of exposure of sugar <u>containing</u> broths to autoclave temperatures does not exceed 45 minutes.
C	1	1.7.12	Media sterility is determined for each load sterilized. Results are recorded

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			and the records maintained.
C	1		1.7.13 Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is made from its individual components.
O	9		1.7.14 Sterile phosphate buffered dilution water is used as the sample diluent.
K	11		1.7.15 The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media			
K	9		1.8.1 Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11		1.8.2 Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13		1.8.3 Stored media are labeled with the storage expiration date or the sterilization date.
K	9		1.8.4 Storage of prepared culture media at room temperature does not exceed <u>seven (7)</u> days.
K	2		1.8.5 Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed <u>one (1)</u> month.
K	11		1.8.6 Storage under refrigeration of prepared culture media with screw-cap closures does not exceed <u>three (3)</u> months.
K	17		1.8.7 All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES			
2.1 Collection and Transportation of Samples			
C	11		2.1.1 Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1		2.1.2 Samples are identified with collector's name, harvest area, sampling station, time and date of collection.
C	9		2.1.3 Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1		2.1.4 A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9		2.1.5 Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.
2.2 Bacteriological Examination of Seawater by the APHA MPN			
C	9		2.2.1 Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
C	2		2.2.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	9, 35 34		2.2.3 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9		2.2.4 In a multiple dilution series not less than <u>three (3)</u> tubes per dilution are used (<u>Five (5)</u> tubes are recommended).

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C	6	2.2.5	In a single dilution series not less than 12 tubes are used (for depuration at least five (5) tubes are used).
C	6	2.2.6	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
K	9	2.2.7	Inoculated tubes are incubated in air at $35 \pm 0.5^\circ\text{C}$.
C	2	2.2.8	Appropriately diluted process control cultures accompany the samples <i>throughout both the presumptive and confirmed phases of incubation</i> . Results are recorded and the records maintained. Positive process control _____ Negative process control _____
K	9	2.2.9	Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
2.3 Confirmed Test for Seawater by APHA MPN			
C	9	2.3.1	Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9	2.3.2	EC medium is used as the confirmatory medium for fecal coliforms.
C	2	2.3.3	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11	2.3.4	Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. (<i>Circle the method of transfer.</i>)
C	9	2.3.5	BGB tubes are incubated at $35 \pm 0.5^\circ\text{C}$.
K	9	2.3.6	BGB tubes are read after 48 ± 3 hours of incubation.
C	9	2.3.7	EC tubes are incubated in a circulating waterbath maintained at $44.5 \pm 0.2^\circ\text{C}$.
C	9	2.3.8	EC tubes are read after 24 ± 2 hours of incubation.
C	9	2.3.9	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.4 Computation of Results – APHA MPN			
K	9	2.4.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7	2.4.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.4.3	Results are reported as MPN/100 mL of sample.
2.5 Bacteriological Examination of Seawater by the MA-1 Method			
C	5	2.5.1	A-1 medium complete is used in the analysis.
C	2, 3 ³⁰	2.5.2	A-1 medium without salicin is used in the analysis. Comparability testing supports use of A-1 medium without salicin. Study records are maintained and are available upon request .
C	5	2.5.3	A-1 medium sterilized for 10 minutes at 121°C .
C	2	2.5.4	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records

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			maintained. Positive productivity control _____ Negative productivity control _____
C	9, 35 34	2.5.5	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in seven (7) seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9	2.5.6	In a multiple dilution series of not less than three (3) tubes per dilution are used (five (5) tubes are recommended).
C	6	2.5.7	In a single dilution series at least 12 tubes are used.
C	6	2.5.8	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	2.5.9	Appropriately diluted process control cultures accompany the samples <i>throughout both resuscitation and waterbath incubation</i> Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	2.5.10	Inoculated tubes are placed in an air incubator at $35 \pm 0.5^\circ\text{C}$ for 3 ± 0.5 hours of resuscitation.
C	5	2.5.11	After 3 ± 0.5 hours resuscitation at 35°C , inoculated tubes are incubated at $44.5 \pm 0.2^\circ\text{C}$ in a circulating waterbath for the remainder of the 24 ± 2 hours.
C	5	2.5.12	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
		2.6 Computation of Results – APHA MPN	
K	9	2.6.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	2.6.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.6.3	Results are reported as MPN/100 mL of sample.
		2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment	
C	23, 24	2.7.1	When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at $44.5 \pm 0.5^\circ\text{C}$ under any loading capacity.
C	23	2.7.2	When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	2.7.3	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	2.7.4	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	2.7.5	Colonies are counted with the aid of magnification.
C	11, 23	2.7.6	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of $0.45 \mu\text{m}$ and certified by the manufacturer for fecal coliform analyses.
C	2	2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
C	2	2.7.8	When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison

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			testing implemented. The results are recorded and this<u>the</u> record is maintained.
K	2, 11		2.7.9 New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2		2.7.10 The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2		2.7.11 Membrane filters which are beyond their expiration date are not used.
O	11		2.7.12 Forceps tips are clean.
O	11		2.7.13 Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.
K	11		2.7.14 Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11		2.7.15 If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or -with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11		2.7.16 Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11		2.7.17 Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26		2.7.18 A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11		2.7.19 The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2		2.7.20 Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
		2.8 Media Preparation and Storage – MF using mTEC Agar	
K	11		2.8.1 Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
C	11		2.8.2 The phosphate buffered saline is properly sterilized.
K	23		2.8.3 A sufficient amount of medium (4-5 mL) is used in each plate.
O	11		2.8.4 Refrigerated prepared plates are stored for no more than <u>two (2)</u> weeks in sealed plastic bags or containers to minimize evaporation.
		2.9 Sample Analyses - MF using mTEC Agar	
C	24		2.9.1 mTEC agar is used.
C	2		2.9.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
C	23, 35 <u>34</u>		2.9.3 The sample is shaken vigorously (25 times in a 12" arc in <u>seven (7)</u> seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration.
C	23		2.9.4 The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25		2.9.5 Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23		2.9.6 Sample volumes are filtered under vacuum.
K	26		2.9.7 The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26		2.9.8 The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.
C	23		2.9.9 The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11		2.9.10 Blanks are run at the beginning of filtration, after every 10th aliquot and at

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			the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).
C	2, 11	2.9.11	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24	2.9.12	Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at $35 \pm 0.5^\circ\text{C}$ for two (2) hours of resuscitation. Alternatively, inoculated plates may be placed in ethafoam prior to air incubation at $44.5 \pm 0.5^\circ\text{C}$ for 24 ± 2 hours.
C	11, 23, 24	2.9.13	After 2 hours of resuscitation at 35°C , the watertight, tightly sealed containers are transferred to a circulating waterbath at $44.5 \pm 0.2^\circ\text{C}$, submerged completely and incubated for 22-24 hours.
2.10 Computation of Results - MF using mTEC Agar			
C	23	2.10.1	All yellow, yellow-green or yellow-brown colonies are counted.
C	23	2.10.2	Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as $>80 \times 100/\text{the volume of sample filtered}$.
C	2, 11, 23	2.10.3	When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11	2.10.4	The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11	2.10.5	Results are reported as CFU/100 mL of sample.
2.11 Bacteriological Analysis of UV Treated Process Water Samples by Membrane Filtration (MF) using mEndo Agar LES – Materials and Equipment			
C	9, 11, 21	2.11.1	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches are used.
C	2	2.11.2	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
C	11, 19, 21	2.11.3	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm diameter with a pore size of $0.45\mu\text{m}$ and certified by the manufacturer for total coliform analysis.
C	2	2.11.4	Membrane filter lot numbers, dates of receipt and expiration dates are recorded and the records maintained.
C	2	2.11.5	If previous lots of agar or membrane filters are not available for comparability testing, an appropriate method for determining lot suitability has been developed and comparison testing implemented when the following has occurred: a. initiating monitoring by mEndo Agar LES; b. changing agar manufacturers; or c. changing brands of membrane filters used. The results are recorded and the records are maintained.
K	2, 11	2.11.6	Recovery of total coliforms from new lots of membrane filters and mEndo Agar LES is compared against the recovery from the previously acceptable lot.
C	2	2.11.7	The sterility of each lot and autoclave batch of membrane filters is verified before use.
K	2	2.11.8	Expired membrane filters are not used.

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K	9, 11, 19, 21	2.11.9	Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
K	11	2.11.10	If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
C	9, 11	2.11.11	Membrane filter assemblies are autoclave sterilized for 15 minutes at 121+/- 2°C prior to the start of a filtration run. A new run occurs when there is a break of 30 minutes or more between the previous filtration run.
O	11, 19, 26, 3635	2.11.12	A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11	2.11.13	The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and the records are maintained.
K	2	2.11.14	Maintenance of the UV sterilization unit is performed as needed. Maintenance is documented and the records maintained.
O	9, 11	2.11.15	Forceps tips are clean and smooth without pitting or corrugations.
2.12 Media Preparation and Storage			
C	9, 11, 19, 21, 3635	2.12.1	mEndo Agar LES is used.
K	11, 21, 3635	2.12.2	mEndo Agar LES is prepared aseptically using pre-sterilized glassware, sterile reagent water and pre-sterilized stir bar.
K	9, 11, 3635	2.12.3	mEndo Agar LES is prepared using 95% alcohol that is not denatured.
C	9, 11, 3635	2.12.4	mEndo Agar LES is brought to near boiling; removed from the heat and tempered at 44-50°C before dispensing.
C	9, 11, 3635	2.12.5	mEndo Agar LES is never autoclaved.
K	9, 11, 3635	2.12.6	A sufficient amount of medium (4-5 mL) is aseptically aliquoted to each culture plate.
O	9, 11, 3635	2.12.7	Prepared plates of mEndo Agar LES are stored at 4°C in the dark for no more than two (2) weeks in sealed plastic bags or containers to minimize evaporation.
C	2	2.12.8	Appropriate, properly diluted positive and negative productivity controls for mEndo Agar LES medium are used. Results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11, 21, 3635	2.12.9	Sterile phosphate buffered water or sterile phosphate buffered saline is used as a sample blank, filter funnel rinse and process and productivity control diluent for UV treated process water samples.
C	11	2.12.10	The phosphate buffered water/saline is properly sterilized and the sterility is tested before being placed in service. Results are recorded and records maintained
2.13 Sample Analysis			
C	9, 11, 3635	2.13.1	The sample is shaken vigorously (25 times in a 12-inch arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration.
C	11, 21, 3635	2.13.2	The membrane filter is placed grid side up within the sterile filter apparatus.
C	11, 26, 3635	2.13.3	A 100 mL quantity of sample is filtered under vacuum.
K	26	2.13.4	The pressure of the vacuum pump does not exceed 15 psi.
C	9, 11, 26,	2.13.5	The sides of the filter funnel are rinsed at least twice with 20-30 mL of

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	3635		sterile phosphate buffered water/saline as appropriate after filtration.
C	9, 11, 3635	2.13.6	The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mEndo Agar LES so that no bubbles form between the filter and the agar.
K	9, 11, 3635	2.13.7	Forceps are dipped in alcohol and flame sterilized between sample filters.
C	11, 3635	2.13.8	Blanks are run at the beginning and at the end of the filtration run to check the sterility of the testing system (phosphate buffered water/saline, filter funnels, forceps, membrane filters, media and culture plates).
C	2, 3635	2.13.9	An appropriate properly diluted positive process control culture accompanies the sample throughout incubation. Results are recorded and the records are maintained.
			Positive process control
C	9, 11, 3635	2.13.10	Inoculated plates are incubated inverted at 35+/- 0.5°C for 22 to 24 hours.
K	2, 9, 11	2.13.11	An appropriate level of humidity is maintained in the incubator to prevent the plates from drying out.
		2.14 Computation of Results	
K	9, 11	2.14.1	Colonies are counted with the aid of magnification.
C	9, 11, 19, 21, 23	2.14.2	All metallic sheen colonies are counted as total coliforms.
C	9, 11, 21, 3635	2.14.3	Results are reported as total coliforms/100mL.
C	11, 20, 3635	2.14.4	When no colonies are observed, results are reported as <1.0 coliform/100mL (nondetectable)
		PART III - SHELLFISH SAMPLES	
		3.1 Collection and Transportation of Samples	
C	9	3.1.1	A representative sample of shellstock is collected.
K	9	3.1.2	Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9	3.1.3	Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9	3.1.4	Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1	3.1.5	Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
		3.2 Preparation of Shellfish for Examination	
K	2, 1 32	3.2.1	Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2, 32	3.2.2	Blades of shucking knives are not corroded.
O	9, 32	3.2.3	The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2, 32	3.2.4	The faucet used for rinsing the shellstock does not contain an aerator.
K	9, 32	3.2.5	Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9, 32	3.2.6	Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	1, 9, 32	3.2.7	Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9, 32	3.2.8	Shellstock are not shucked directly through the hinge.

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C	9, 32		3.2.9	Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	2, 9		3.2.10	At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.
K	9		3.2.11	A representative sample of at least 12 shellfish is used for the analysis.
K	2		3.2.12	The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
O	9		3.2.13	Sterile phosphate buffered dilution water is used as the sample diluent.
C	9		3.2.14	Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9		3.2.15	APHA <i>Recommended Procedures for the Examination of Sea Water And Shellfish</i> , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA				
C	9		3.3.1	Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. <i>(Circle the medium used.)</i>
C	2		3.3.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9		3.3.3	Immediately (within <u>two (2)</u> minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9		3.3.4	No fewer than <u>five (5)</u> tubes per dilution are used in a multiple dilution MPN series.
C	9		3.3.5	Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6		3.3.6	In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2		3.3.7	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9		3.3.8	Inoculated media are incubated at $35 \pm 0.5^{\circ}\text{C}$.
K	10		3.3.9	Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA				
C	9		3.4.1	EC medium is used as the confirmatory medium.
C	2		3.4.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11		3.4.3	Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. <i>(Circle the method of transfer.)</i>
C	9		3.4.4	EC tubes are incubated in a circulating waterbath at $44.5 \pm 0.2^{\circ}\text{C}$
K	9		3.4.5	EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9		3.4.6	The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.

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3.5 Computation of Results for MPN Analyses			
K	9		3.5.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.
K	7		3.5.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9		3.5.3 Results are reported as MPN/100 grams of sample.
3.6 Standard Plate Count Method			
O	20		3.6.1 A standard plate count (SPC) analysis may be performed in conjunction with the analysis for fecal coliform organisms.
K	9		3.6.2 In the standard plate count procedure at least four <u>(4)</u> plates are used, duplicates of two <u>(2)</u> dilutions. One <u>(1)</u> of the dilutions should produce colonies of 30 to 300 per plate.
K	2		3.6.3 Fifteen <u>15</u> to 20 mL of tempered sterile plate count agar is used per plate.
C	9		3.6.4 Agar tempering bath maintains the agar at 44-46°C.
C	9		3.6.5 An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9		3.6.6 Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in <u>seven</u> <u>(7)</u> seconds) before plating.
C	9		3.6.7 Not more than 1 mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11		3.6.8 Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9, 21		3.6.9 Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four <u>(4)</u> high.
K	9		3.6.10 Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1		3.6.11 A hand tally or its equivalent is used for accuracy in counting.
3.7 Computation of Results -SPC			
K	9		3.7.1 Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19		3.7.2 Colony counts are reported as CFU/<u>grams</u> of sample.
3.8 Bacteriological Analysis of Shellfish Using the ETCP			
C	2, 3		3.8.1 Prepared modified MacConkey agar is used on the day that it is made.
K	3		3.8.2 Double strength modified MacConkey agar is used.
C	3		3.8.3 Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3		3.8.4 Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50°C until used.
K	2, 3		3.8.5 Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3		3.8.6 The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9		3.8.7 The sample homogenate is cultured within <u>two</u> <u>(2)</u> minutes of blending.
C	2, 3		3.8.8 Six <u>(6)</u> grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.
K	3		3.8.9 Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2, 3, 22		3.8.10 The container is gently swirled or slowly inverted once to mix the contents,

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			which are subsequently distributed uniformly over six (6) plates.
C	1		3.8.11 Media and diluent sterility are determined with each use. Results are recorded and the records maintained.
C	1		3.8.12 Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture _____ Negative control culture _____
C	3, 13		3.8.13 When solidified, the plates are placed inverted into an air incubator at $-45.5 \pm 0.5^{\circ}\text{C}$ for 18 to 30 hours of incubation.
C	2		3.8.14 Plates are stacked no more than three (3) high in the incubator.
C	2		3.8.15 Appropriately diluted pour plated process control cultures accompany each set of samples throughout incubation. The results are recorded and the records maintained. Positive process control _____ Negative process control _____
3.9 Computation of Results - ETCP			
K	11		3.9.1 Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1		3.9.2 A hand tally or its equivalent is used to aid in counting.
C	3, 6		3.9.3 All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3		3.9.4 Results are reported as CFU/100 grams of sample.
Bacteriological Examination of Soft-shelled Clams and American Oysters Shellfish Meats for Male Specific Coliphage (MSC)			
3.10 MSC Equipment and Supplies			
K	302		3.10.1 Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold <u>at least</u> 100—125 mL.
€	27, 28		3.10.2 The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	92		3.10.3 2 The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
€	27, 28		3.10.4 Sterile 0.22 µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1		3.10.5 3 The sterility of each batch/lot of pre-sterilized <u>or reusable</u> syringes, <u>and syringe filters, and/or filter units</u> is determined. Results are recorded and records maintained.
K	4		3.10.6 The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28 <u>2</u>		3.10.7 4 The balance used provides a sensitivity of at least mg (0.01g.).
C	27, 28, 31		3.10.8 5 The temperature of the incubator used is maintained at $36 \pm 1^{\circ}\text{C}$.
K	2		3.10.6 The temperature of the freezer is maintained at $\leq -15^{\circ}\text{C}$.
C	28 <u>1</u>		3.10.9 7 <u>Sterile</u> The sterility of disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
3.11 MSC Media Preparation			
K	28, 31		3.11.1 Media preparation and sterilization is according to the validated method.
K	27, 28, 31		3.11.2 Bottom agar, double strength soft agar and growth broth are prepared from their individual components Antibiotic solutions are filter sterilized using sterile 0.22 µm pore size filters.
K	27, 28		3.11.3 Soft agar is prepared double strength in volumes of 2.5 mL.
€	27, 28		3.11.4 The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.

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O	27, 28, <u>31</u>	3.11.5— <u>3</u> Storage of the bottom agar under refrigeration does not exceed 1 month <u>six (6) weeks</u> .
K	<u>27, 28</u> 2	3.11.6— <u>4</u> Unsterilized soft agar is stored at $\leq -20^{\circ}\text{C}$ - 15°C for up to <u>three (3) months</u> .
K	27, 28, <u>31</u>	3.11.7— <u>5</u> The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.
K	<u>27, 28</u>	3.11.8 Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	<u>28, 31</u>	3.11.6 Storage under refrigeration of prepared growth broth with screw-cap closures shall not exceed three (3) months and with loose fitting closures shall not exceed one (1) month.
K	<u>2, 27, 28, 31</u>	3.11.9— <u>7</u> Bottom agar plates <u>stored under refrigeration</u> are allowed to reach room temperature before use.
3.12 Preparation of Host Culture for MSC Analysis		
C	<u>28, 31</u>	3.12.1 E. coli Famp ATCC 700891 is the bacterial host strain.
K	<u>27, 28, 31</u>	3.12.2 Host cell growth broth is tempered at $36 \pm 1^{\circ}\text{C}$ prior to inoculation with host cells.
K	<u>27, 28, 31</u>	3.12.3 Several host cell colonies are transferred to a tube of tempered growth broth and incubated at $36 \pm 1^{\circ}\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	<u>27, 28, 31</u>	3.12.4 After inoculation, the host cell growth broth culture is not shaken.
3.12-13 Preparation of the Soft-Shelled Clams and American Oysters Shellfish for MSC Analysis		
K	<u>2, 4 36</u>	3. 12 <u>13</u> .1 Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2	3. 12 <u>13</u> .2 The blades of shucking knives are not corroded.
O	9	3. 12 <u>13</u> .3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	3. 12 <u>13</u> .4 The faucet used for rinsing the shellfish does not contain an aerator.
K	9	3. 12 <u>13</u> .5 The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	3. 12 <u>13</u> .6 The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9	3. 12 <u>13</u> .7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9	3. 12 <u>13</u> .8 Shellfish are not shucked through the hinge.
C	9	3. 12 <u>13</u> .9 The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	3. 12 <u>13</u> .10 A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19	3. 12 <u>13</u> .11 The sample is weighed to the nearest 0.1 gram.
C	<u>28, 31</u>	3.13.2 Two (2) times the weight of the sample of sterile growth broth, by volume, is added.
C	<u>28, 31</u>	3.13.13 Samples are blended at high speed for 180 seconds.
3.13-14 MSC Sample Analysis		
C	<u>28</u>	3.13.1 E. coli Famp ATCC 700891 is the bacterial host strain used in this procedure.
K	<u>27, 28</u>	3.13.2 Host cell growth broth is tempered at $36 \pm 1^{\circ}\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	<u>27, 28</u>	3.13.3 Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^{\circ}\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	<u>27, 28</u>	3.13.4 After inoculation, the host cell growth broth culture is not shaken.
C	<u>28</u>	3.13.5 A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting

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			the MSC.
C	28		3.13.6 The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28		3.13.7 The elution mixture is homogenized at high speed for 180 seconds.
C	28, 31		3.13.8 14.1 Immediately after blending, 33 grams of the homogenized homogenate elution mixture are is weighed into <u>a</u> centrifuge tubes.
C	28, 31		3.13.9 14.2 The homogenized elution mixture is centrifuged for 15 minutes at 9000 x g at 4°C.
C	27, 28, 31		3.13.10 14.3 The supernatant is pipetted off transferred to a new sterile tube , weighed, and the weight recorded.
C	27, 28, 31		3.13.11 14.4 The supernatant is allowed to warm to room temperature about 20 to 30 minutes prior to analysis .
K	27, 28, 31		3.13.12 14.5 The autoclaved soft agar is tempered and held at 51 ± 1°C throughout the period of sample analysis.
K	27, 28, 31		3.13.13 14.6 Two hundred 200 microliters (0.2 mL) of log phase host strain <i>E coli</i> is added to the tempering tempered soft agar immediately prior to adding the sample supernatant.
K	27, 28, 31		3.13.14 14.7 The sample supernatant is shaken or vortexed before being added to the tempering tempered soft agar.
C	27, 28, 31		3.13.15 14.8 2.5 mL of sample supernatant (avoiding bubbles where necessary) is added to each a tube of tempering tempered soft agar.
C	27, 28, 31		3.13.16 14.9 The tube of soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28, 31		3.13.17 14.10 The soft agar/sample supernatant/host cell mixture tube contents is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.
C	28, 31		3.13.18 14.11 Ten (10) plates are used for analysis of each sample with , 2.5 mL of sample supernatant per plate for a total of 25 mL of supernatant analyzed per sample, unless fewer than 25 mL of supernatant is obtained from the sample in which all supernatant is plated.
K	27, 28, 31		3.13.19 14.12 Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are recorded and records maintained. Positive control _____
K	27, 28, 31		3.13.20 14.13 Room temperature g Growth broth is used as the negative control or blank.
K	27, 28, 31		3.13.21 14.14 Type strain MS2 (ATCC 15597- B1) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2		3.13.22 14.15 A negative control plate is plated at the beginning and end of each set of samples analyzed.
K	27, 28, 31		3.13.23 14.16 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28, 31		3.13.24 14.17 All plates are incubated at 36 ± 1° C for 18 ± 2 hours.
			3.14-15 Computation of Results - MSC
C	27		3.14 15.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 31, 32, 36		3.14 15.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten plates, the count reported value is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/ 100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as > 20,000 PFU/100 grams.
K	28, 31		3.14 15.3 The formula used for determining the density of MSC in PFU/100 grams is: (0.364) (N) (Ws), where N = total number of plaques counted on all 10 plates and Ws = weight of the supernatant used.
O	92		3.14 15.4 The MSC count is rounded off conventionally to give a whole number.

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LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
MICROBIOLOGICAL COMPONENT: (Part I-III)	
A. Results	
Total # of Critical (C) Nonconformities in Parts I-III	_____
Total # of Key (K) Nonconformities in Parts I-III	_____
Total # of Critical, Key and Other (O) Nonconformities in Parts I-III	_____
B. Criteria for Determining Laboratory Status of the Microbiological Component:	
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <ul style="list-style-type: none"> a. The total # of Critical nonconformities is ≥ 4 or b. The total # of Key nonconformities is ≥ 13 or c. The total # of Critical, Key and Other is ≥ 18 <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but ≤ 3.</p>	
C. Laboratory Status (circle appropriate)	
Does Not Conform	Provisionally Conforms
Conforms	
<p>Acknowledgment by Laboratory Director/Supervisor:</p> <p>All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____.</p> <p>Laboratory Signature: _____ Date: _____</p> <p>LEO Signature: _____ Date: _____</p>	

Receptor Binding Assay for Paralytic Shellfish Poisoning (PSP)		
PART I – Quality Assurance		
ITEM		
CODE	REF	
1.1 Quality Assurance (QA) Plan		
K	1, 2, 3	1.1.1 Written Plan (Check <input checked="" type="checkbox"/> those items which apply).
		a. Organization of the Laboratory.
		b. Staff training requirements. Training must include radiation lab safety.
		c. Standard operating procedures (SOPs).
		d. Internal quality control measures for equipment, their calibration maintenance, repair, performance and rejection criteria established.
		e. Laboratory safety. Radiation safety practices (e.g., handling and disposal) must be included.
		f. Internal performance assessment.
		g. External performance assessment.
C	2	1.1.2 The QA plan is implemented.
1.2 Educational/Experience Requirements		
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analysts meet the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree in microbiology, biology or other appropriate discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial laboratories, the analysts must have at least a high school diploma and at least three months of experience in laboratory sciences.
C	6	1.2.5 Training regarding radiation laboratory safety, handling and disposal practices and verification of licensing must be provided is documented and records are maintained.
C	15	1.2.6 Laboratory has a Nuclear Regulatory Commission (NRC) or equivalent state license for the use of tritiated saxitoxin in this assay. Alternatively, the laboratory uses less than 50 µCi per year and adheres to the American Radiolabeled Chemical (ARC) exemption status.
1.3 Work Area		
O	2	1.3.1 The work area is adequate for the workload and storage.
K	2	1.3.2 The work area is clean and well lighted.
K	2	1.3.3 The work area has adequate temperature control.
O	3	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
C	3,4	1.3.5 The work area is located in an appropriate space designated for low-level radiation work. Radioactive materials are only handled and manipulated in designated areas which are clearly identified and labeled accordingly.
1.4 Laboratory Equipment		
C	4	1.4.1 Any lab equipment that may come into contact with [³ H]-STX at any point in the preparation or assay procedures must be specially labelled and must

Laboratory Evaluation Checklist – Receptor Binding Assay for Paralytic Shellfish Poisoning (PSP)

			remain in the work area designated for low-level radiation work.
O	5		1.4.2 The pH meter has a standard accuracy of 0.1 pH units.
K	7		1.4.3 The pH electrodes being used consist of a pH half cell and reference half cell or equivalent combination electrode/triode free from silver/silver chloride (Ag/AgCl) or contains an ion exchange barrier to prevent the passage of silver (Ag) ions into the substance being measured.
K	3, 8		1.4.4 The pH meter is calibrated daily when in use. Results are recorded and records maintained.
K	1		1.4.5 The effect of temperature on the pH has been compensated for by an ATC probe, use of a triode, or by manual adjustment.
K	1		1.4.6 The pH meter manufacturer instructions are followed for calibration, or a minimum of two (2) standard buffer solutions is used to calibrate the pH meter. If the calibration sequence of standard buffer solutions is not stipulated by the manufacturer, the first must be near the isopotential point (pH 7) and the second near the expected sample (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	9		1.4.7 Electrode acceptability is determined daily or with each use by the millivolt procedure or through determination of the slope.
K	6		1.4.8 pH paper in the appropriate pH range (i.e., 1-5), if used, measures accurately to a minimum of 0.5 pH units over the covered pH range.
K	6		1.4.9 The differing sensitivities in weight measurements required by the various steps in the assay are met by the balance(s) being used. a. To prepare Phenyl methylsulfonyl fluoride solution (PMSF), the balance used must have a sensitivity of at least 0.001 gram at a load of 1 gram. b. For sample extraction, the balance used must have a sensitivity of at least 0.1 gram at a load of 100 grams. c. For MOPS buffer preparation, the balance used must have a sensitivity of at least 0.01 gram at a load of 100 grams.
K	1, 3		1.4.10 Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use.
			1.4.11 Balances must be calibrated by an external service at least once per year. Results are recorded and records maintained.
K	2		1.4.12 Refrigerator temperatures are maintained between 0 and 4 °C. Freezer security for ³ HSTX and cold STX must meet state and federal requirements for these materials.
K	1		1.4.13 Refrigerator temperatures are monitored at least once daily on workdays. Results are recorded and records maintained.
C	4, 6, 10		1.4.14 Freezer temperature used to store [³H] STX standard, rat brain membrane tissue preparation, interassay calibration standard (QC check) and archived shellfish tissue homogenate is maintained at -80 °C or below. Freezer security for ³HSTX and cold STX must meet state and federal requirements for these materials.
K	6, 10		1.4.15 Freezer temperature used for all other purposes is maintained at -20 °C or below.
O	1		1.4.16 Freezer temperature is monitored at least once daily on workdays. Results are recorded and records maintained.
O	8		1.4.17 All glassware is clean.
C	3		1.4.18 An alkaline or acid-based detergent is used for washing glassware/labware.
C	1		1.4.19 With each load of labware/glassware washed, the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali as appropriate) with aqueous 0.04% bromothymol blue (BTB) solution. Results are recorded and records maintained.
C	6		1.4.20 Micropipettors are calibrated for the appropriate volumes used and checked

Laboratory Evaluation Checklist – Receptor Binding Assay for Paralytic Shellfish Poisoning (PSP)

			annually for accuracy. Results are recorded and records are maintained.
C	11		1.4.21 Scintillation counter is serviced according to manufacturer specifications and calibrated annually. Results are recorded and records maintained.
C	4		1.4.22 Minimum radiation safety equipment and protocols include the following: A wipe-test is conducted in the radiation work area as described in the QA plan. Results are recorded and records maintained.
1.5 Reference Solution Reagent Storage, Preparation and Security			
C	12		1.5.1 [³H] STX standard is stored in a freezer at -80 °C or below.
C	10		1.5.2 Concentration of [³H] STX standard is calculated from the lot information provided by the supplier with each batch.
K	6		1.5.3 Unopened diHCl STX standard may be stored at room temperature or refrigerated.
C	10		1.5.4 Preparation of MOPS assay buffer includes the following: a. 100 mM MOPS/L. b. 100 mM choline chloride/L. c. pH adjustment to 7.4 with NaOH. e. refrigerated storage at 4 °C. d. Maintained ice cold while in use.
C	10		1.5.6 Bulk standard curve dilutions are stored at 4 °C for up to one (1) month.
K	1		1.5.7 Reagent water is distilled or deionized (<i>circle appropriate choice</i>) and is analyzed monthly for the following criteria, with all results recorded and records maintained: a. Exceeds 0.5 megohm-cm resistivity (2 megohm-cm in-line) or less than 2.0 μSiemens/cm conductivity at 25 °C (<i>circle appropriate choice</i>). b. Residual chlorine is at a non-detectable level (<0.1 ppm). Specify method of determination _____. c. Water contains <100 CFU/mL using the heterotrophic plate count method.
1.6 Rat Brain Membrane Tissue Preparation and Storage			
C	10		1.6.1 MOPS/choline chloride/phenyl methylsulfonyl fluoride (PMSF), pH 7.4 is used in preparing rat brain membrane tissue. PMSF is added to MOPS/choline chloride fresh on the day of use.
C	10		1.6.2 The cerebral cortex of 6-week old Sprague-Dawley rats is used in membrane tissue preparations, placed in iced MOPS/choline chloride/PMSF buffer (pH 7.4; 1 brain/12.5 mL) and homogenized with no visible chunks remaining in the homogenate. This procedure is repeated until twenty (20) rat brains have been processed.
C	10		1.6.3 The homogenized cerebral cortex tissue from the twenty (20) rat brain cortices is pooled and centrifuged at 20000 x g for 15 minutes at 4 °C.
K	10		1.6.4 The pellet of the centrifuged rat brain tissue preparation is fully resuspended in ice cold MOPS/choline chloride/PMSF buffer (up to 10 mL/brain).
K	10		1.6.5 The resuspended rat brain tissue preparations are pooled and the centrifuge tubes used for these preparations are rinsed with a small amount of MOPS/choline chloride/PMSF buffer to recover all the rat brain tissue.
K	10		1.6.6 The total volume of the pooled rat brain tissue is adjusted to 200 mL with MOPS/choline chloride/PMSF buffer while iced.
K	10		1.6.7 The iced contents of the pooled rat brain tissue are blended using a Polytron at 70% power or a small hand- held blender at low speed for 20 seconds to obtain a homogeneous membrane tissue preparation.
C	10		1.6.8 Two (2) mL/tube of the pooled, homogeneous rat brain membrane tissue preparation is aliquoted into cryovials, frozen and stored at -80 °C for up to six (6) months.
1.7 Rat Brain Membrane Tissue Protein Receptor Determination			
C	10		1.7.1 The protein/receptor concentration of the rat brain membrane tissue preparation is determined for each new batch using a Pierce Micro BCA Protein Assay Reagent Kit No. 23235 (micro plate method) or No. 23225 (tube

Laboratory Evaluation Checklist – Receptor Binding Assay for Paralytic Shellfish Poisoning (PSP)

			method) or equivalent.
C	10		1.7.2 The dilution of the protein/receptor concentration of the rat brain membrane tissue preparation needed to obtain a working stock of 1 mg/mL is determined.
K	10		1.7.3 Dilutions of the protein/receptor concentration of the rat brain membrane tissue preparation of less than 1:4 are not used as they may be too viscous.
PART II – Analysis of Shellfish Samples for PSP Toxins – RBA			
2.1 Collection and Transportation of Samples			
C	5		2.1.1 A representative sample of shellfish is collected.
K	5		2.1.2 Shellfish samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	5		2.1.3 Shellfish samples are labeled with the collector’s name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	5		2.1.4 Immediately after collection, shellstock samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10 °C with ice or cold packs for transport to the laboratory.
K	6, 13		2.1.5 Time from collection to initiation of the extraction should not exceed 24 hours. However, if significant delays are anticipated or if they occur, the laboratory has an appropriate contingency plan in place to handle these samples. For samples shipped live in accordance with 2.1.4, the contingency plan ensures samples remain within allowable temperature tolerances and animals are alive upon receipt. The contingency plan also addresses field and/or laboratory processing that ensures the integrity of the sample or extract until initiation of the assay. For example, samples are washed, shucked, drained and processed as follows: a. refrigerated or frozen until extracted; b. homogenized and frozen until extracted; or c. extracted, the supernatant decanted, and refrigerated or frozen until assayed.
2.2 Preparation of Samples for Analysis – Homogenization			
C	5, 6		2.2.1 At least 12 animals are used per sample, or the laboratory has an appropriate contingency plan for dealing with non-typical species of shellfish or collection conditions.
O	5		2.2.2 The outside of the shell is thoroughly cleaned with fresh water.
O	5		2.2.3 Shellstock are opened by cutting the adductor muscles.
O	5		2.2.4 The inside surfaces of the shells and meats are rinsed with fresh water to remove sand or other foreign material.
O	5		2.2.5 Shellfish meats are removed from the shell by separating the adductor muscles and tissue connecting at the hinge.
C	5		2.2.6 Damage to the body of the mollusk is minimized in the process of opening.
O	5		2.2.7 Shucked shellfish are drained on a #10 mesh sieve or equivalent without layering for 5 minutes.
K	5		2.2.8 Pieces of shell and drainage are discarded.

C	5, 6	<p>2.2.4 Meats are blended at high speed until homogenous (60 – 120 seconds), using the following criteria:</p> <ul style="list-style-type: none"> a. Freshly drained/air dried meats are placed into the blender for homogenization. b. Previously frozen shucked, rinsed, and drained meats are completely thawed, then placed in the blender <u>with all freeze-thaw liquid</u> for homogenization. c. Previously frozen homogenates are completely thawed then placed in the blender <u>with all freeze-thaw liquid</u> for homogenization.
K	6, 13	2.2.5 Homogenates should be extracted immediately. If homogenates must be stored, they should be frozen.
2.3 Preparation of Samples for Analysis – Extraction		
K	5, 10	2.3.1 0.1 M HCl is used for extractions.
K	5, 10	2.3.2 <u>At least five (5) grams of tissue +/- 0.1g is extracted using an equal amount a 1:1 mass to volume ratio of 0.1 M HCl.</u>
C	10	2.3.3 The pH of the sample is checked and adjusted as necessary to between 3.0–4.0.
C	10	2.3.4 Adjustment of the pH is accomplished by dropwise addition of either 5 N HCl or 0.1 N NaOH, as appropriate, while constantly stirring the sample.
C	6	2.3.5 The sample is promptly brought to a boil at 99.0 +/- 1.0 °C and gently boiled for 5 minutes.
O	6	2.3.6 The sample is boiled under adequate ventilation (e.g., fume hood).
O	10	2.3.7 The sample is allowed to cool to room temperature.
C	10	2.3.8 The pH of the cooled mixture after boiling is between 3.0 - 4.0, adjusted if necessary, with the dropwise addition of 5 M HCl to lower the pH or 0.1 M NaOH to raise the pH, as appropriate, while constantly stirring the mixture.
K	5, 10	2.3.9 The volume of the sample is adjusted to the original (pre-boiling) volume, by adding 0.001N HCl (pH 3 water).
K	10	2.3.10 The sample is stirred gently to homogeneity, then treated as follows: <ul style="list-style-type: none"> a. The sample is allowed to settle to remove particulates, then the supernatant is carefully decanted into a clean container; then b. an aliquot of the sample is centrifuged at 3000 x g for 10 minutes, then the supernatant is carefully decanted into a clean container.
K	6, 10	2.3.11 The sample extract is analyzed immediately, refrigerated at 4 °C in a sealed container for up to 24 hours, or frozen at -20 °C.
2.4 Sample Assay		
K	6	2.4.1 One analyst performs the entire plate set-up for the assay.
K	6	2.4.2 Microtubes containing dilutions and samples are vortexed immediately before dispensing.
K	10	2.4.3 The standard curve consists of at least 7 concentrations (minimum 6×10^{-10} M and maximum 6×10^{-6} M).
C	10	2.4.4 The rat brain membrane tissue preparation is kept on ice and mixed often during addition to the plate to maintain a homogenous suspension.
K	10	2.4.5 Each day an assay is conducted, a standard curve, reference blank, and an inter-assay QC calibration standard is required. However, filter plates of the same lot must be used if the assay requires multiple plates to accommodate all samples. If the filter plate lot changes over the course of a day, a new standard curve must be performed for the new lot of filter plates. <u>An inter-assay QC calibration and reference blank are required for each plate analyzed.</u>
C	10	2.4.6 The standard curve, reference blank, interassay QC calibration standard, and test samples are all run in triplicate.
K	10	2.4.7 Assay buffer is added to the plate before any other components of the assay, in order to properly wet the filter membrane.

Laboratory Evaluation Checklist – Receptor Binding Assay for Paralytic Shellfish Poisoning (PSP)

K	10	2.4.8 All wells of the plate (including any unused wells) are filled with MOPS/choline chloride buffer during vacuum filtration, in order to ensure even pressure and filtration across the plate.
C	10	2.4.9 Appropriate scintillation cocktail is used, depending on the type of scintillation counter (traditional or microplate).
K	10	2.4.10 If ³ H STX working solution is checked for counts per minute (CPM) it should be and is consistent and within 15% of the expected value.
C	10	2.4.11 An appropriate dark adaptation interval is employed, based on type of scintillation counter (traditional or microplate).
K	10	2.4.12 Standard curve fitting is calculated using appropriate software program.
C	10	2.4.13 Slope of standard curve is between -0.8 and -1.2 (the theoretical slope is -1.0). If the slope falls outside these criteria, the assay results are rejected and the assay must be repeated.
C	10	2.4.14 The relative standard deviation of triplicate CPM for standards and samples must be less than 30%. If greater than 30%, the assay results are rejected and the assay must be repeated.
C	10	2.4.15 The IC₅₀ is in acceptable range (2.0 nM +/- 30%). If the IC₅₀ is outside this range, the assay results are rejected and the assay must be repeated
C	10	2.4.16 The inter-assay QC calibration standard (QC check) sample is in the acceptable range (3 nM +/- 30%). If the QC check sample is outside this range, the assay results are rejected and the assay must be repeated.
C	10	2.4.17 Sample dilutions are quantified only if B/B₀ is between 0.2 – 0.7. If B/B₀ is greater than 0.7, then the sample is reported as below the limit of detection. If B/B₀ is less than 0.2, then the sample should be further diluted and repeated if a quantification is needed.
K	4	2.4.18 Assay materials are cleaned and disposed of in accordance with federal, state, and local requirements.
2.5 Calculation of Sample Toxicity		
C	10	2.5.1 When more than one dilution falls within B/B₀ of 0.2 – 0.7, all wells corresponding to these dilutions are used to calculate sample toxicity.
C	10	<p>2.5.2 Sample toxicity is calculated as follows:</p> <p style="text-align: center;">(nM STX equiv.) x (sample dilution) x (210 µL total volume/35 µL sample = mM STX equivalent in extract</p> <p style="text-align: center;">(nM STX diHCl equiv. in extract) x 1L/1000 mL x 372 ng/nmol x 1 µg/1000 ng =µg STX diHCl equiv./mL</p> <p style="text-align: center;">µg STX diHCl equiv./mL x mL extract/g shellfish x 1000 g/kg =µg STX diHCl equiv./kg</p>
C	14	2.5.3 Any value equal to or greater than 80 µg STX diHCl equiv./100 g) of sample is actionable.
C		Shellfish Program Management is made aware of positive result. Laboratory action to identify positive result is: _____.

References:

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5. American Public Health Association (APHA). 1970. Recommended Procedures for the Examination of Sea Water and Shellfish, Fourth Edition. APHA, Washington, D.C.
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9. Consult pH electrode product literature.
10. Association of Official Analytical Chemists (AOAC). 2016. Official Method 2011.27 Paralytic Shellfish Toxins (PSTs) in Shellfish Receptor Binding Assay.
11. Consult instrument manufacturer instructions.
12. Technical Data Sheet, American Radiolabeled Chemicals, Inc. 101 Arc Drive, St. Louis, MO 63146.
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14. U. S. Food and Drug Administration (FDA) and Interstate Shellfish Sanitation Conference (ISSC). 2017. *NSSP Guide for the Control of Molluscan Shellfish*. FDA/ISSC, Washington D.C. and Columbia, S.C.
15. U. S. Nuclear Regulatory Commission Materials, Section 30.18, 10 CFR Part 30, and American Radiolabeled Chemicals Licenses.

<p>PUBLIC HEALTH SERVICE U.S. FOOD AND DRUG ADMINISTRATION OFFICE OF FOOD SAFETY SHELLFISH AND AQUACULTURE POLICY BRANCH 5001 CAMPUS DRIVE COLLEGE PARK, MD 20740-3835 TEL. 240-402-2151/2055/4960 FAX 301-436-2601 CESANDSSLEOS@FDA.HHS.GOV</p>		
<p>SHELLFISH LABORATORY EVALUATION CHECKLIST Domoic Acid (Amnesic Shellfish Poisoning; ASP) HPLC-UV</p>		
LABORATORY:		
ADDRESS:		
TELEPHONE:	FAX:	EMAIL:
DATE OF EVALUATION:	DATE OF REPORT:	LAST EVALUATION:
LABORATORY REPRESENTED BY:		TITLE:
LABORATORY EVALUATION OFFICER:		SHELLFISH SPECIALIST:
		REGION:
OTHER OFFICIALS PRESENT:		TITLE:
Items which do not conform are noted by:		
<p>C – Critical K - Key O - Other NA - Not Applicable Conformity is noted by a “1”</p>		

PART I – QUALITY ASSURANCE		
Code	REF	Item Description
1.1 Quality Assurance (QA) Plan		
K	5, 8	1.1.1 Written Plan adequately covers all the following: (check '1' those that apply) a. Organization of the laboratory. b. Staff training requirements. c. Standard operating procedures. d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance and rejection criteria established. e. Laboratory safety. f. Internal performance assessment.
C	5	1.1.2 QA Plan is implemented.
1.2 Educational/Experience Requirements		
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial/private laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, chemistry, or another appropriate discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial/private laboratories, the analyst must have at least a high school diploma and shall have at least three months of experience in laboratory sciences.
C	5	1.2.5 LC-Operator must be trained in the operation and maintenance of a basic liquid chromatography system.
1.3 Work Area		
O	5, 8	1.3.1 Adequate for workload and storage.
O	8	1.3.2 Clean and well lighted.
O	8	1.3.3 Adequate temperature control.
O	8	1.3.4 All work surfaces are nonporous and easily cleaned.
1.4 Laboratory Equipment		
K	6	1.4.1 The pH meter has a standard accuracy of 0.1 unit. [Only applicable if using the sample cleanup procedure]
K	5	1.4.2 The pH meter is calibrated daily when in use. Results are recorded and records are maintained. [Only applicable if using the sample cleanup procedure]
K	8	1.4.3 Effect of temperature has been compensated for by an ATC probe, use of a triode or by manual adjustment. [Only applicable if using the sample cleanup procedure]

K	8	1.4.4 The pH meter manufacturer instructions are followed for calibration or a minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second must be near the expected sample pH (i.e., pH 2, 4 or 11) as appropriate. Standard buffer solutions are used once and discarded. [Only applicable if using the sample cleanup procedure]
K	5, 11	1.4.5 Electrode acceptability is determined daily or with each use following either slope or millivolt procedure. [Only applicable if using the sample cleanup procedure]
K	6, 2	1.4.6 The balances being used provide an appropriate sensitivity at the weights of use, at least 0.1 g for laboratory precision balances and 0.1 mg for analytical balances.
K	8	1.4.7 The balance calibration is checked monthly using NIST class S, ASTM class 1 or 2 weights or equivalent. Results are recorded and records are maintained.
K	1	1.4.8 Refrigerator temperature is maintained between 0 and 4-6 °C.
K	8	1.4.9 Refrigerator temperature is monitored at least once daily. Results are recorded and records maintained.
K	4,15	1.4.10 Freezer temperature is maintained at -10 °C or below.
K	8	1.4.11 Freezer temperature is monitored at least once daily. Results are recorded and records maintained.
C	13	1.4.12 All in-service thermometers are properly calibrated and immersed.
K	5	1.4.13 All glassware is clean.
K	4	1.4.14 A high performance liquid chromatography system (HPLC) equipped with the following is used: <ul style="list-style-type: none"> a. mobile phase system delivering a pulse-free flow of 1.0 mL/min, b. solvent degasser, c. autosampler (refrigerated preferred) with loop suitable for 20 µL injections, d. temperature controlled column compartment capable of controlling temperature at 40 °C, e. ultraviolet detector/diode array detector able to achieve the required sensitivity at a wavelength (A) of 242 nm, and f. a data collection system (e.g., computer, integrator).
K	2	1.4.15 Autopipettors are calibrated for the appropriate volumes used and checked annually for accuracy. Results are recorded and records are maintained.
K	4	1.4.16 A solid phase extraction (SPE) vacuum manifold capable of holding 3 mL cartridges is used. [Only applicable if using the sample cleanup procedure]
O	4	1.4.17 A centrifuge capable of holding 50 mL polypropylene tubes is used.
1.5 Reagents and Reference Solution Preparation and Storage		
C	4, 15	1.5.1 All solvents and reagents used are analytical or LC grade materials.
O	8	1.5.2 Water contains < 100 CFU/ml as determined monthly using the heterotrophic plate count method. Results are recorded and records are maintained. (Not required for bottled reagent grade or HPLC grade water when used immediately upon opening. If the bottle of water is not used entirely immediately, the water must be tested as above prior to continued use.)

K	8	1.5.3 Reagents are properly stored and labeled with the date of receipt, date opened or date prepared and expiration date.
C	4, 15	1.5.4 The mobile phase system used to analyze domoic acid consists of: 10% aqueous acetonitrile (v/v) and 0.1% trifluoroacetic acid (TFA).
O	4	1.5.5 Mobile phase is filtered before use if the HPLC does not have a degasser.
C	7	1.5.6 Only certified reference materials are used for standard solutions. Source of the reference standard: _____
K	4, 15	1.5.7 A cartridge wash solution is made up of 1 volume acetonitrile to 9 volumes of water (i.e., 10% aqueous acetonitrile). [Only applicable if using the sample cleanup procedure]
K	4	1.5.8 Citrate buffer (0.5 M, pH 3.2) is made up by dissolving 40.4 g citric acid monohydrate and 14 g triammonium citrate in 400 mL water, then adding 50 mL acetonitrile and diluting the total to 500 mL with water [or equivalent buffer]. [Only applicable if using the sample cleanup procedure]
C	7	1.5.9 NRC CRM Zero-Mus or a negative control is used as a blank to ensure that there is no carry over between samples/standards. Source of the negative control: _____
C	7	1.5.10 All primary standards are stored appropriately as per supplier recommendations.
C	7	1.5.11 All standards used are within expiration date.
C	2	1.5.12 All standards are prepared either gravimetrically or using positive displacement pipettes.
C	4, 15	1.5.13 Working standards are made up from primary standard by dilution with the toxin-free, extraction solvent (i.e., 50% aqueous methanol). Dilution with toxin-free, cartridge wash solution (aqueous acetonitrile) is allowed if using the diluted crude sample or the sample cleanup procedure.
C	7	1.5.14 Zero-Mus is stored according to manufacturer's instructions.
C	2	1.5.15 Quality Control shellfish tissues are stored frozen.
1.6 Collection and Transportation of Samples		
O	6, 1	1.6.1 Shellstock are collected in clean, waterproof, puncture resistant containers.
K	6, 1	1.6.2 Samples are appropriately labeled with the collector's name, type of shellstock, the harvest area, and time and date of collection.
C	6, 1	1.6.3 Immediately after collection, shellstock samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10 °C with ice or cold packs for transport to the laboratory.

K	14, 2	1.6.4 Time from collection to initiation of the extraction should not exceed 24 hours. However, if significant delays are anticipated or if they occur, the laboratory has an appropriate contingency plan in place to handle the samples. For samples shipped live in accordance with 1.6.3, the contingency plan ensures samples remain within allowable temperature tolerances and animals are alive upon receipt. The contingency plan also addresses field and/or laboratory processing that ensures the integrity of the sample or extract until initiation of the assay. For example, samples are washed, shucked, drained and processed as follows: a. refrigerated or frozen until extracted; b. homogenized and frozen until extracted; or c. extracted, the supernatant decanted, and refrigerated or frozen until assayed.
C	2	1.6.5 Frozen shucked product or homogenates are allowed to thaw completely and all liquid is included as part of the sample before being processed further.
PART II – EXAMINATION OF SHELLFISH FOR ASP TOXINS		
2.1 Preparation of Sample		
C	6, 1	2.1.1 At least 12 animals are used per sample or the laboratory has an appropriate contingency plan for dealing with non-typical species of shellfish (e.g., 3 geoduck gut balls).
O	6	2.1.2 The outside of the shell is thoroughly cleaned with fresh water.
O	6	2.1.3 Shellstock are opened by cutting the adductor muscles.
O	6	2.1.4 The inside surfaces of the shells are rinsed with fresh water to remove sand and other foreign materials.
O	6	2.1.5 Shellfish meats are removed from the shell by separating the adductor muscles and tissue connecting at the hinge.
C	6	2.1.6 Damage to the body of the mollusk is minimized in the process of opening.
O	6	2.1.7 Shucked shellfish are drained on a #10 mesh sieve or equivalent without layering for 5 minutes.
K	6	2.1.8 Pieces of shell and drainage are discarded.
C	2, 6	2.1.9 Drained meats or previously cooled/refrigerated shucked meats and their drip loss liquid or thawed homogenates with their freeze-thaw liquid are blended at high speed until homogenous (60-120 seconds).
2.2 Sample Extraction		
K	4,6	2.2.1 Sample homogenates are extracted as soon as possible (preferably the same day) or stored in the freezer at -10 °C or below.
C	4	2.2.2 Four (4) grams of homogenized sample is weighed into a 50 mL polypropylene centrifuge tube and subsequently extracted.
C	4	2.2.3 The sample homogenate is extracted with 16 mL extraction solvent (1:1 methanol:water [also referred to as 50% aqueous methanol]).
K	4, 15	2.2.4 Homogenate/extract mixture is centrifuged and filtered before analysis.
K	4, 15	2.2.5 The filtered extract is injected into the HPLC or loaded into the autosampler immediately.

K	4		2.2.6 When crude samples are diluted, dilutions are made by diluting 1 mL of filtered sample supernatant into a 5 mL volumetric flask and diluted with water to 5 mL.
K	4, 15		2.2.7 Crude extracts are sealed tightly and stored at -10 °C or below.
2.3 Sample Cleanup (Optional)			
O	4, 15		2.3.1 Three (3) mL SAX cartridges (500 mg silica derivatized with quaternary ammonium silane) are used for cleanup.
K	4		2.3.2 The SAX cartridge is conditioned with 6 mL methanol, followed by 3 mL water, followed by 3 mL extraction solvent (1:1 methanol:water).
C	4, 15		2.3.3 The cartridge is not allowed to run dry during conditioning through sample loading.
K	4, 15		2.3.4 Five (5) mL of filtered extract is loaded onto the cartridge and flowed slowly (~1 drop/s) until sample meniscus reaches the top of cartridge packing, discarding effluent.
K	4, 15		2.3.5 Five (5) mL of wash solution (1:9 acetonitrile:water) is loaded to the cartridge and washed slowly (~1 drop/s) until meniscus reaches the top of cartridge packing, discarding effluent.
K	4		2.3.6 0.5 mL of citrate buffer (0.5 M, pH 3.2) is loaded to the cartridge and flowed slowly (~1 drop/s) until meniscus reaches the top of cartridge packing, discarding effluent.
K	4, 15		2.3.7 A 2 mL volumetric tube is placed under the cartridge and any domoic acid is eluted into the tube by loading and flowing as much citrate buffer as needed slowly (~ 1 drop/s) until the 2 mL mark is reached on the tube.
C	4, 15		2.3.8 The solution is thoroughly mixed before withdrawing an aliquot for analysis.
K	4, 15		2.3.9 The cleaned up extract is injected into the HPLC or loaded into the autosampler immediately.
2.4 Analysis			
C	2		2.4.1 A standard calibration curve (of at least six concentrations) is performed daily. Results are recorded and records are maintained.
K	4, 15		2.4.2 Twenty (20) µL of extract is injected for analysis.
K	2		2.4.3 Samples are stored in the sample compartment of the autosampler at 4 °C during analysis. Otherwise samples must be analyzed within 9 hours if the autosampler is held at room temperature.
K	4, 15		2.4.4 A column heater is used and the temperature is maintained at 40 °C during the analysis.
C	4		2.4.5 The appropriate analytical column is used: 25 cm x 4.6 mm id packed with 5 µm Vydac 201TP octadecylsilica or equivalent.
K	2		2.4.6 The column is stored following the manufacturer's instructions when not in use.
O	2		2.4.7 If a precolumn in-line filter and/or a compatible guard column (e.g., 201GCC54T) are/is used, rejection criteria are established to determine when to change the filter/guard column.

C	2	2.4.8 Dead volume in the system is minimized by the use of short lengths of connecting tubing of small internal diameter between the sample injector and the column and between the column and detector.
2.5 System Suitability		
C	2	2.5.1 The correlation coefficient for the linear regression of the calibration standards must be ≥ 0.990 for domoic acid.
C	3	2.5.2 The resolution and retention time criteria must ensure complete baseline resolution of L-tryptophan and domoic acid.
K	2	2.5.3 Peak asymmetry is routinely monitored to evaluate the performance of the column. Results are recorded and records maintained.
C	2	2.5.4 The column is replaced when a measure of peak asymmetry becomes $<0.90.5$ or >1.3 .
C	2,4	2.5.5 Daily injection schedules must include the adequate frequency of injection standards and extraction blanks based on an assessment of individual standard toxin variability and lack of carry over.
C	2	2.5.6 Repeated injections of calibrated standards/samples agree within $\pm 5\%$ (as determined through the use of the coefficient of variation).
2.6 Calculation of Sample Toxicity		
C	4, 15	<p>2.6.1 The toxicity of the individual toxins is calculated as follows:</p> $\mu\text{g/g domoic acid (DA)} = \text{DA injected} \times \frac{V}{W} \times (F)$ <p>where:</p> <p>DA injected = the concentration in $\mu\text{g/ml}$ of the extract injected; V = total volume of homogenate and extraction solvent (mL); W = weight (g) of tissue homogenate extracted (e.g., 4 g); and F = dilution factor (e.g., if SAX cleanup or crude sample dilution are performed).</p> <p>The concentration of DA injected may be determined using the nearest standard or the equation of the day's standard curve.</p>
C	4, 15	2.6.2 Calculated domoic acid concentrations include the sum of domoic acid and epimer peaks, when the epimer represents 5% or more of the peak area.
C	12	2.6.3 Any value at or above 20 ppm (mg/kg or $\mu\text{g/g}$) domoic acid is actionable.
REFERENCES		
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3. AOAC Official Method 991.26 Domoic Acid in Mussels. Liquid Chromatography Method. First Action 1991. Final Action 1999.		
4. Quilliam, M.A., M. Xie, and W.R. Hardstaff. 1995. J. AOAC Int. 78(2): 543-554.		

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National Shellfish Sanitation Program (NSSP) Guide for the Control of Molluscan Shellfish: 2017 Revision

LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
AMNESIC SHELLFISH POISON (ASP or domoic acid) COMPONENT: PARTS I AND II	
A. Results Total # of Critical (C) Nonconformities _____ Total # of Key (K) Nonconformities _____ Total # of Critical, Key, and Other (O) Nonconformities _____	_____ _____ _____
B. Criteria for Determining Laboratory Status of the ASP (domoic acid) Component <ol style="list-style-type: none"> 1. Conforms Status: The ASP component of this Laboratory is in conformity with NSSP requirements if all of the following apply. <ol style="list-style-type: none"> a. No Critical nonconformities. b. and <6 Key nonconformities. c. and <12 Total nonconformities. 2. Provisionally Conforms Status: The ASP component of this laboratory is determined to be provisionally conforming to NSSP requirements if all of the following apply. <ol style="list-style-type: none"> a. the number of critical nonconformities is ≥ 1 but < 4. b. and < 6 Key nonconformities. c. and < 12 Total nonconformities. 3. Does Not Conform Status: The ASP component of this laboratory is not in conformity with NSSP requirements when any of the following apply. <ol style="list-style-type: none"> a. The total # of Critical nonconformities is ≥ 4. b. or the total # of Key nonconformities is ≥ 6. c. or the total # of Critical, Key, or Other is ≥ 12. 	
C. Laboratory Status (circle appropriate) Does Not Conform – Provisionally Conforms – Conforms	

National Shellfish Sanitation Program (NSSP) Guide for the Control of Molluscan Shellfish: 2017 Revision

Acknowledgement by Laboratory Director/Supervisor:	
All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before_____.	
Laboratory Signature:_____	Date:_____
LEO Signature:_____	Date:_____

PUBLIC HEALTH SERVICE U.S. FOOD AND DRUG ADMINISTRATION OFFICE OF FOOD SAFETY SHELLFISH AND AQUACULTURE POLICY BRANCH 5100 PAINT BRANCH PARKWAY COLLEGE PARK, MD 20740-3835 TEL. 240-402-2151/2055/4960 FAX 301-436-2601		
SHELLFISH LABORATORY EVALUATION CHECKLIST		
LABORATORY:		
ADDRESS:		
TELEPHONE:	FAX:	EMAIL:
DATE OF EVALUATION:	DATE OF REPORT:	LAST EVALUATION:
LABORATORY REPRESENTED BY:		TITLE:
LABORATORY EVALUATION OFFICER:		SHELLFISH SPECIALIST:
		REGION:
OTHER OFFICIALS PRESENT:		TITLE:
Items which do not conform are noted by:		
C – Critical K - Key O - Other NA - Not Applicable Conformity is noted by a “√”		

PART I – QUALITY ASSURANCE		
Code	REF	Item Description
1.1 Quality Assurance (QA) Plan		
K	5, 8	1.1.1 Written Plan adequately covers all the following: (check \checkmark those that apply) a. Organization of the laboratory. b. Staff training requirements. c. Standard operating procedures. d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance and rejection criteria established. e. Laboratory safety. f. Internal performance assessment. g. External performance assessment.
C	5	1.1.2 QA Plan is implemented.
1.2 Educational/Experience Requirements		
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial/private laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, chemistry, or another appropriate discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial/private laboratories, the analyst must have at least a high school diploma and shall have at least three months of experience in laboratory sciences.
C	5	1.2.5 LC-Operator must be competent in the operation and maintenance of a basic liquid chromatography system.
1.3 Work Area		
O	5, 8	1.3.1 Adequate for workload and storage.
O	8	1.3.2 Clean and well lighted.
O	8	1.3.3 Adequate temperature control.
O	8	1.3.4 All work surfaces are nonporous and easily cleaned.
1.4 Laboratory Equipment.		
O	6	1.4.1 The pH meter has a standard accuracy of 0.1 unit.
K	6	1.4.2 pH paper in the appropriate range (i.e. 1-4), if used, is used with minimum accuracy of 0.5 pH units.
K	10	1.4.3 pH electrodes consist of pH half-cell and reference half-cell or equivalent combination electrode/triode (free from Ag/AgCl or contains an ion exchange barrier to prevent passage of Ag ions into the medium that may result in inaccurate pH readings).

K	5	1.4.4 pH meter is calibrated daily when in use. Results are recorded and records are maintained.
K	8	1.4.5 Effect of temperature has been compensated for by an ATC probe, use of a triode or by manual adjustment.
K	8	1.4.6 A minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second must be near the expected sample pH (i.e. pH 2, 4 or 11) as appropriate. Standard buffer solutions are used once and discarded.
K	5, 11	1.4.7 Electrode acceptability is determined daily or with each use following either slope or millivolt procedure.
K	6	1.4.8 The balances being used provide an appropriate sensitivity at the weights of use, at least 0.1 g for laboratory precision balances and 0.1 mg for analytical balances.
K	8, 9	1.4.9 The balance calibration is checked monthly using NIST class S, ASTM class 1 or 2 weights or equivalent. Results are recorded and records are maintained.
K	1	1.4.10 Refrigerator temperature is maintained between 0 and 4 °C.
K	8	1.4.11 Refrigerator temperature is monitored at least once daily. Results are recorded and records maintained.
K	1	1.4.12 Freezer temperature is maintained at -20 °C or below.
K	8	1.4.13 Freezer temperature is monitored at least once daily. Results are recorded and records maintained.
C	13	1.4.14 All in-service thermometers are properly calibrated and immersed.
K	5	1.4.15 All glassware is clean.
K	3	1.4.16 A high performance liquid chromatography system (HPLC) equipped with the following is used: <ul style="list-style-type: none"> a. binary mobile phase system delivering a pulse-free flow of 0.5-2.0 mL/min, b. solvent degasser, c. autosampler (refrigerated preferred) with loop suitable for 5-30 µL injections, d. temperature controlled column compartment capable of controlling temperature between 10 – 50 °C, and e. fluorescence detector able to achieve the required sensitivity at an excitation wavelength (λ) of 330 nm and emission of 390 nm.
K	3, 4	1.4.17 The post-column reaction system used is equipped with the following: <ul style="list-style-type: none"> a. reactor module capable of maintaining 85 °C, b. dual reagent pumps capable of delivering accurate flows of 0.4 mL/min, and c. if applicable, a reaction coil (knitted or equivalent) having a total volume of 1 mL and a length of 5 m x 0.5 mm.
K	6	1.4.18 Autopipettors are calibrated for the appropriate volumes used and checked annually for accuracy. Results are recorded and records are maintained.
K	3	1.4.19 A boiling water bath with sufficient volume to cover the sample/acid mixture is used for extraction.
O	3	1.4.20 Centrifuge capable of holding 50 mL polypropylene tubes.
K	3	1.4.21 Microcentrifuge capable of holding 1.5 mL microcentrifuge tubes and generating a minimum of 16000 g or equivalent is used.
1.5 Reagents and Reference Solution Preparation and Storage		
C	3	1.5.1 All solvents and reagents used are analytical or LC grade materials.
C	8	1.5.2 Water is glass distilled or deionized and exceeds 0.5 megaohm resistance or is less than 2 µSiemens/cm conductivity at 25 °C to be tested and recorded monthly for resistance or conductivity and the results are recorded.
K	8	1.5.3 Water is analyzed for residual chlorine monthly and is at a nondetectable level

		(≤0.1 ppm) Results are recorded and records are maintained.
K	8	1.5.4 Water contains < 100 CFU/ml as determined monthly using the heterotrophic plate count method. Results are recorded and records are maintained.
K	8	1.5.5 Reagents are properly stored and labeled with the date of receipt, date opened or date prepared and expiration date.
C	3	1.5.6 The binary mobile phase system used to analyze the GTX and STX toxins consists of: 1.5.6.1 Mobile Phase A, which contains 11 mM heptane sulfonate and 5.5 mM phosphoric acid (H ₃ PO ₄), pH 7.1. 1.5.6.2 Mobile Phase B, which contains 11 mM heptane sulfonate, 16.5 mM H ₃ PO ₄ and 11.5% acetonitrile (MeCN), pH 7.1.
C	3	1.5.7 The binary mobile phase system used to analyze the C toxins consists of 1.5.7.1 Mobile Phase A, which contains 2 mM tetrabutyl ammonium phosphate, pH 5.8. 1.5.7.2 Mobile Phase B, which contains 2 mM tetrabutyl ammonium phosphate in 4% acetonitrile, pH 5.8.
C	3	1.5.8 The post-column oxidant consists of 100 mM H ₃ PO ₄ and 5 mM periodic acid (H ₅ IO ₆), pH 7.8.
C	3	1.5.9 The post-column acid used is 0.75 M nitric acid (HNO ₃).
C	3	1.5.10 The heptane sulfonate used in mobile phase A and mobile phase B to analyze for GTX and STX toxins is prepared the day of use or refrigerated for up to one week.
C	3	1.5.11 The pH of mobile phases and the post-column oxidant are adjusted as follows: a. Mobile phase A and mobile phase B for the GTX and STX toxins are adjusted to 7.1 with ammonium hydroxide (NH ₄ OH), b. Mobile phase A and mobile phase B for the C toxins are adjusted to 5.8 in one direction only with 10% acetic acid (HOAc) if too basic or 1% NH ₄ OH if too acidic, and c. The post-column oxidant is adjusted to 7.8 with 5 M sodium hydroxide (NaOH).
O	3	d. Mobile phases and post-column reagents are filtered before use if the HPLC does not have a degreaser.
C	3,7	1.5.12 Only certified reference materials are used for standard solutions. Source of the reference standard:
C	7	1.5.13 NRC Zero-Mus or a negative control matched matrix is used as a matrix blank as appropriate. Source of the negative matrix:
C	7	1.5.14 All primary standards are stored appropriately as per supplier recommendations.
C	7	1.5.15 All standards used are within expiration date.
C	3	1.5.16 All standards are prepared gravimetrically.
K	3	1.5.17 Intermediate mixes of primary standards are made up in 0.003 M HCl for the GTX/STX toxins or pH 5 glass distilled/deionized water for the C toxins labeled with the date of preparation and the expiration date and stored appropriately. The pH of the glass distilled/deionized water is adjusted when necessary by the dropwise addition of 10% acetic acid (HOAc).
C	3	1.5.18 Working standards are made up from primary standard or intermediate mixes by dilution with toxin-free, deproteinated, matrix matched extracts.
C	7	1.5.19 Zero-Mus is stored according to manufacturer's instructions.

C	2	1.5.20 Quality Control shellfish tissues are stored frozen.
C	7	1.5.21 Working standards are labeled with the date of preparation, stored appropriately and used within 3 months of preparation.
1.6 Collection and Transportation of Samples		
O	6	1.6.1 Shellstock are collected in clean, waterproof, puncture resistant containers.
K	6	1.6.2 Samples are appropriately labeled with the collector's name, type of shellstock, the harvest area, and time and date of collection.
C	6	1.6.3 Immediately after collection, shellstock samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10 °C with ice or cold packs for transport to the laboratory.
K	14	1.6.4 Time from collection to initiation of the extraction should not exceed 24 hours. However, if significant delays are anticipated or if they occur, the laboratory has an appropriate contingency plan in place to handle the samples. For samples shipped live in accordance with 1.6.3, the contingency plan ensures samples remain within allowable temperature tolerances and animals are alive upon receipt. The contingency plan also addresses field and/or laboratory processing that ensures the integrity of the sample or extract until initiation of the assay. For example, samples are washed, shucked, drained and processed as follows: a. refrigerated or frozen until extracted; b. homogenized and frozen until extracted; or c. extracted, the supernatant decanted, and refrigerated or frozen until assayed.
C	6	1.6.5 Frozen shucked product or homogenates are allowed to thaw completely and all liquid is included as part of the sample before being processed further.
PART II – EXAMINATION OF SHELLFISH FOR PSP TOXINS		
2.1 Preparation of Sample		
C	6	2.1.1 At least 12 animals are used per sample or the laboratory has an appropriate contingency plan for dealing with non-typical species of shellfish.
O	6	2.1.2 The outside of the shell is thoroughly cleaned with fresh water.
O	6	2.1.3 Shellstock are opened by cutting the adductor muscles.
O	6	2.1.4 The inside surfaces of the shells are rinsed with fresh water to remove sand and other foreign materials.
O	6	2.1.5 Shellfish meats are removed from the shell by separating the adductor muscles and tissue connecting at the hinge.
C	6	2.1.6 Damage to the body of the mollusk is minimized in the process of opening.
O	6	2.1.7 Shucked shellfish are drained on a #10 mesh sieve or equivalent without layering for 5 minutes.
K	6	2.1.8 Pieces of shell and drainage are discarded.
C	6	2.1.9 Drained meats or previously cooled/refrigerated shucked meats and their drip loss liquid or thawed homogenates with their freeze-thaw liquid are blended at high speed until homogenous (60-120 seconds).
2.2 Digestion of Sample		
K	6	2.2.1 Sample homogenates are extracted as soon as possible (preferably the same day) or stored in the freezer.
C	3	2.2.2 Five (5) grams of homogenized sample is weighed into a 50 mL polypropylene centrifuge tube and subsequently extracted.
K	3	2.2.3 The sample homogenate is extracted in a 1:1 w/v ratio with 0.1 M HCl.

K	3	2.2.4 Homogenate/acid mixture is vortexed thoroughly before boiling to completely mix the contents.
C	3	2.2.5 To prevent toxin transformation, the pH of the homogenate/acid mixture before boiling is 3.0 ± 1.0, adjusted if necessary with the dropwise addition of either 5 M HCl to lower the pH or 0.1 M NaOH to raise the pH.
C	3	2.2.6 Samples in capped 50 mL polypropylene centrifuge tubes are extracted in a boiling water bath for 5 minutes.
K	3	2.2.7 The pH of the cooled mixture after boiling is 3.0 ± 1.0 , adjusted if necessary with the dropwise addition of 5 M HCl. Any sample with a pH of less than 2.0 is discarded and extracted again.
K	3	2.2.8 The homogenate/acid mixture is allowed to separate by gravity or by centrifugation.
2.3 Deproteination		
C	3	2.3.1 500-1000 μL of sample extract is deproteinated with 25-50 μL of 30% trichloroacetic acid, vortexed thoroughly and centrifuged at $\sim 16,000 g$ for 5 minutes.
C	3	2.3.2 The pH of the deproteinated extract is adjusted with 35 μL of 12.0 M NaOH vortexed thoroughly and centrifuged at $\sim 16,000 g$ for 5 minutes.
K	3	2.3.3 An aliquot of the deproteinated supernatant is filtered through a 0.2 μm filter.
2.4 Analysis		
C	2	2.4.1 A standard calibration curve (of at least six concentrations) is performed upon initial instrument set up, following any major hardware maintenance activity, or when the continuing calibration verification (CCV) indicates significant drift ($> 30\%$ for individual toxin) from the calibration. Results are recorded and records are maintained.
K	3	2.4.2 10 μL is injected for GTX/STX toxins and 5 μL is injected for C-toxins.
K	3	2.4.3 Samples are stored in the sample compartment of the autosampler at 4 °C during analysis. Otherwise samples must be analyzed within 20 hours if the autosampler is held at room temperature.
K	3	2.4.4 A column heater that is capable of maintaining 30-40 °C for the GTX/STX toxins and 10-20 °C for the C toxins is used in the analysis.
C	3	2.4.5 The appropriate analytical column is used. a. GTX/STX Toxins: Agilent Zorbax Bonus-RP column, 4.6 mm x 150 mm, 3.5 μm or equivalent. b. C Toxins: Thermo BetaBasic 8, 4.6 mm x 250 mm, 5 μm or equivalent.
2.5 System Suitability		
K	2	2.5.1 The correlation coefficient for the linear regression of the calibration standards must be ≥ 0.990 for each individual toxin.

C	3	<p>2.5.2 The resolution and retention time criteria that must be met are:</p> <ul style="list-style-type: none"> a. For GTX and STX toxins, the matrix peak must be at least 70% baseline resolved between GTX3 and GTX2. b. For GTX and STX toxins, GTX5 must be at least 40% baseline resolved between dcGTX3 and dcGTX2. c. For GTX and STX toxins, dcSTX and STX must be at least 70% baseline resolved. d. For GTX and STX toxins, the retention time of GTX4 must be between 5 and 7 minutes. e. For the C toxins, C2 must be at least 70% baseline resolved between C1 and C2. f. For the C toxins, the retention time of C1 must be between 4 and 7 minutes. 																																
C	2	<p>2.5.3 Daily injection schedules must include the adequate frequency of injection standards based on an assessment of individual standard toxin variability. Variability in peak response must be less than 10% for calculation of toxicity in samples.</p>																																
2.6 Calculation of Toxicity																																		
C	4	<p>2.6.1 The toxicity of the individual toxins is calculated as follows:</p> $\mu\text{gSTXdiHCleq}/100\text{g} = \mu\text{M} \times \frac{372.2}{1000\text{mL}} \times \frac{\text{Fvol}}{\text{Ext.vol}} \times \left(\frac{\text{Wt} + \text{Vol}}{\text{Wt}} \right) \times \text{ReTx} \times 100$ <p>14. Where:</p> <ul style="list-style-type: none"> μM = Concentration of toxin in the extract, in μM; Fvol = Final volume of the deproteinized extract (e.g. 560 μL); Ext.vol = Volume of crude extract used (e.g. 500 μL); Wt = Weight of sample used; Vol = Volume of acid extractant used (e.g. 5 mL); and ReTx = Relative toxicity of toxin vs. Saxitoxin. <p style="text-align: center;">Relative Toxicity Values</p> <table border="1" style="margin-left: auto; margin-right: auto;"> <thead> <tr> <th>Toxin</th> <th>ReTx</th> <th>Toxin</th> <th>ReTx</th> </tr> </thead> <tbody> <tr> <td>GTX1</td> <td>0.9940</td> <td>NEO</td> <td>0.9243</td> </tr> <tr> <td>GTX2</td> <td>0.3592</td> <td>STX</td> <td>1.0000</td> </tr> <tr> <td>GTX3</td> <td>0.6379</td> <td>dcSTX</td> <td>0.5131</td> </tr> <tr> <td>GTX4</td> <td>0.7261</td> <td>C1</td> <td>0.0060</td> </tr> <tr> <td>GTX5</td> <td>0.0644</td> <td>C2</td> <td>0.0963</td> </tr> <tr> <td>dcGTX2</td> <td>0.1538</td> <td>C3</td> <td>0.0133</td> </tr> <tr> <td>dcGTX3</td> <td>0.3766</td> <td>C4</td> <td>0.0576</td> </tr> </tbody> </table> <p>15.</p>	Toxin	ReTx	Toxin	ReTx	GTX1	0.9940	NEO	0.9243	GTX2	0.3592	STX	1.0000	GTX3	0.6379	dcSTX	0.5131	GTX4	0.7261	C1	0.0060	GTX5	0.0644	C2	0.0963	dcGTX2	0.1538	C3	0.0133	dcGTX3	0.3766	C4	0.0576
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C	3	<p>2.6.2 The individual toxicities for each toxin are summed to obtain the overall sample toxicity in μg STX equivalents/100 g ($\mu\text{g}/100\text{g}$).</p>																																
C	12	<p>2.6.3 Any value at or above 80 μg STX equivalents /100 g of meat is actionable.</p>																																

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LABORATORY	DATE			
LABORATORY REPRESENTATIVE:				
PARALYTIC SHELLFISHPOISON COMPONENT: PARTS I AND II				
A. Results Total# of Critical (C) Nonconformities Total# of Key (K) Nonconformities Total# of Critical, Key, and Other (O) Nonconformities	_____ _____ _____			
B. Criteria for Determining Laboratory Status of the PSP, PCOX Component 1. Conforms Status: The PSP, PCOX component of this Laboratory is in conformity with NSSP requirements if a ll of the following apply. a. No Critical nonconformities. b. and <6 Key nonconformities. c. and <12 Total nonconformities. 2. Provisionally Conforms Status: The PSP, PCOX component of this laboratory is determined to be provisionally conforming to NSSP requirements if all of the following apply. a. the number of critical nonconformities is ≥ 1 but < 4. b. and <6 Key nonconformities. c. and <12 Total nonconformities. 3. Does Not Conform Status : The PSP, PCOX component of this laboratory is not in conformity with NSSP requirements when any of the following apply. a. The total# of Critical nonconformities is ≥ 4 . b. or the total# of Key nonconformities is ≥ 6 . c. or the total# of Critical, Key, or Other is ≥ 12 .				
C. Laboratory Status (circle appropriate) <table style="width: 100%; border: none;"> <tr> <td style="width: 33%; text-align: center;">Does Not Conform</td> <td style="width: 33%; text-align: center;">Provisionally Conforms</td> <td style="width: 33%; text-align: center;">Conforms</td> </tr> </table>		Does Not Conform	Provisionally Conforms	Conforms
Does Not Conform	Provisionally Conforms	Conforms		
Acknowledgement by Laboratory Director/Supervisor: All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____. Laboratory Signature: _____ Date: _____ LEO Signature: _____ Date: _____				

PART 1 - QUALITY ASSURANCE		
CODE	REF.	ITEM
K	8, 11	1.1 Quality Assurance (QA) Plan
		1.1.1 Written Plan (Check those items which apply.)
		a. Organization of the laboratory.
		b. Staff training requirements.
		c. Standard operating procedures.
		d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		e. Laboratory safety.
		f. Internal performance assessment.
		g. External performance assessment.
C	8	1.1.2 QA Plan Implemented.
K	11	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s)_____
		1.2 Educational/Experience Requirements
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three months of experience in laboratory science
		1.3 Work Area
O	8,11	1.3.1 Adequate for workload and storage.
K	11	1.3.2 Clean, well-lighted.
K	11	1.3.3 Adequate temperature control.
O	11	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
		1.4 Laboratory Equipment
O	9	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	1.4.4 pH meter is calibrated daily or with each use Results are recorded and records maintained.
K	11	1.4.5 A minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	8,15	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt procedure or through determination of the slope. (<i>Circle the method used.</i>)

K	9		1.4.7 Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13		1.4.8 Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11		1.4.9 Refrigerator temperature(s) are monitored at least once daily on workdays Results are recorded and records maintained.
K	1		1.4.10 Refrigerator temperature is maintained at 0 to 4°C.
C	9		1.4.11 The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11		1.4.12 Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9		1.4.13 Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11		1.4.14 Temperature of the waterbath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9		1.4.15 The thermometers used in the waterbath are graduated in at least 0.1°C increments.
C	13		1.4.16 The waterbath has adequate capacity for workload.
K	9		1.4.17 The level of water in the waterbath covers the level of liquid in the incubating tubes.
K	8, 11		1.4.18 Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4		1.4.19 All working thermometers are appropriately immersed.
C	29		1.4.20 Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	11		1.4.21 A standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9		1.4.22 Standards thermometers are checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination_____.
C	29		1.4.23 Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of ≤ ±0.05°C are used as the laboratory standards thermometer. (Circle the thermometer type used.)
K	13		1.4.24 The accuracy of working thermometers is checked annually against the standards thermometer either at the temperatures at which they are used or by ice point determination. Results are recorded and records maintained.
O	11		1.4.25 Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing			
O	9		1.5.1 Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding materials.
K	9		1.5.2 Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.
K	9		1.5.3 Sample containers are made of glass or some other inert material.
O	9		1.5.4 Dilution bottles and tubes are made of borosilicate glass or plastic and closed with rubber stoppers, caps or screw caps with nontoxic liners.

K	9		1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9		1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1mL used to deliver 0.1 mL aliquots.
K	9		1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9		1.5.8	In washing reusable pipettes, a succession of at least three fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2		1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11		1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination				
K	9		1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8		1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	30, 33, 34		1.6.3	The autoclave provides sterilization conditions suitable to the load contents. Sterilization temperature range may be 119°C - 124°C as determined by the lab's equipment Quality Assurance Verification Testing and recommended practices from the media manufacturer. Sterilization is determined for each load using a working maximum registering thermometer or an appropriate working temperature monitoring device.
K	11		1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16		1.6.5	The autoclave standards thermometer is checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory; or, is checked in-house at the steam point (100°C) if it has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination _____
K	1		1.6.6	Working autoclave thermometers are checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11		1.6.7	Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11		1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13		1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11		1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180°C.
K	9		1.6.11	A thermometer capable of determining temperatures accurately in the range of 160 to 180°C is used to monitor the operation of the hot-air sterilizing oven
K	13		1.6.12	Records of temperatures and exposure times are maintained for the operation of the hot-air sterilizing oven during use.

K	11		1.6.13 Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Records are maintained.
K	11		1.6.14 Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C.
C	1		1.6.15 The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1		1.6.16 The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9		1.6.17 Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9		1.6.18 Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2 hours.
C	2		1.6.19 The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2		1.6.20 The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18		1.6.21 Hardwood applicator transfer sticks are properly sterilized. Method of sterilization _____
C	2		1.6.22 The sterility of the hardwood applicator transfer sticks is checked routinely. Results are recorded and the records maintained.
O	13		1.6.23 Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation			
K	3, 5		1.7.1 Media is commercially dehydrated except in the case of medium A-1 which must be prepared from the individual components and modified MacConkey agar which may be prepared from its components.
K	11		1.7.2 Media is prepared according to manufacturer's instructions.
O	11		1.7.3 Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11		1.7.4 Dehydrated media are labeled with date of receipt and date opened.
C	12		1.7.5 Caked or expired media or media components are discarded.
C	11		1.7.6 Reagent water is distilled or deionized (circle appropriate choice), tested monthly and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) or is less than 2.0 µSiemens/cm conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11		1.7.7 Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level (<0.1 mg/L). Results are recorded and the records maintained. Specify method of determination_____.
K	11		1.7.8 Reagent water contains<100 CFU/mL as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11		1.7.9 Media prepared from commercial dehydrated components are sterilized according to the manufacturer's instructions.
K	9		1.7.10 The volume and concentration of media in the tube are suitable for the amount of sample inoculated.
C	11		1.7.11 Total time of exposure of sugar broths to autoclave temperatures does not exceed 45 minutes.
C	1		1.7.12 Media sterility is determined for each load sterilized. Results are recorded and the records maintained.

C	1		1.7.13 Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is made from its individual components.
O	9		1.7.14 Sterile phosphate buffered dilution water is used as the sample diluent.
K	11		1.7.15 The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media			
K	9		1.8.1 Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11		1.8.2 Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13		1.8.3 Stored media are labeled with the storage expiration date or the sterilization date.
K	9		1.8.4 Storage of prepared culture media at room temperature does not exceed 7 days.
K	2		1.8.5 Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed 1 month.
K	11		1.8.6 Storage under refrigeration of prepared culture media with screw-cap closures does not exceed 3 months.
K	17		1.8.7 All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES			
2.1 Collection and Transportation of Samples			
C	11		2.1.1 Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1		2.1.2 Samples are identified with collector's name, harvest area, sampling station, time and date of collection.
C	9		2.1.3 Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1		2.1.4 A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9		2.1.5 Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.
2.2 Bacteriological Examination of Seawater by the APHA MPN			
C	9		2.2.1 Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
C	2		2.2.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control
C	9, 35		2.2.3 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9		2.2.4 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6		2.2.5 In a single dilution series not less than 12 tubes are used (for depuration at least 5 tubes are used).

C	6	2.2.6	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
K	9	2.2.7	Inoculated tubes are incubated in air at $35 \pm 0.5^\circ\text{C}$.
C	2	2.2.8	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
K	9	2.2.9	Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
2.3 Confirmed Test for Seawater by APHA MPN			
C	9	2.3.1	Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9	2.3.2	EC medium is used as the confirmatory medium for fecal coliforms.
C	2	2.3.3	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11	2.3.4	Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. (<i>Circle the method of transfer.</i>)
C	9	2.3.5	BGB tubes are incubated at $35 \pm 0.5^\circ\text{C}$.
K	9	2.3.6	BGB tubes are read after 48 ± 3 hours of incubation.
C	9	2.3.7	EC tubes are incubated in a circulating waterbath maintained at $44.5 \pm 0.2^\circ\text{C}$.
C	9	2.3.8	EC tubes are read after 24 ± 2 hours of incubation.
C	9	2.3.9	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.4 Computation of Results – APHA MPN			
K	9	2.4.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7	2.4.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.4.3	Results are reported as MPN/100 mL of sample.
2.5 Bacteriological Examination of Seawater by the MA-1 Method			
C	5	2.5.1	A-1 medium complete is used in the analysis.
C	2, 31	2.5.2	A-1 medium without salicin is used in the analysis. Comparability testing supports use of A-1 medium without salicin. Study records are available.
C	5	2.5.3	A-1 medium sterilized for 10 minutes at 121°C .
C	2	2.5.4	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____

C	9, 35	2.5.5	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9	2.5.6	In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	2.5.7	In a single dilution series at least 12 tubes are used.
C	6	2.5.8	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	2.5.9	Appropriately diluted process control cultures accompany the samples <i>throughout both resuscitation and waterbath incubation</i> Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	2.5.10	Inoculated tubes are placed in an air incubator at $35 \pm 0.5^\circ\text{C}$ for 3 ± 0.5 hours of resuscitation.
C	5	2.5.11	After 3 ± 0.5 hours resuscitation at 35°C , inoculated tubes are incubated at $44.5 \pm 0.2^\circ\text{C}$ in a circulating waterbath for the remainder of the 24 ± 2 hours.
C	5	2.5.12	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.6 Computation of Results – APHA MPN			
K	9	2.6.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	2.6.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.6.3	Results are reported as MPN/100 mL of sample.
2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment			
C	23, 24	2.7.1	When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at $44.5 \pm 0.5^\circ\text{C}$ under any loading capacity.
C	23	2.7.2	When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	2.7.3	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	2.7.4	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	2.7.5	Colonies are counted with the aid of magnification.
C	11, 23	2.7.6	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of $0.45 \mu\text{m}$ and certified by the manufacturer for fecal coliform analyses.
C	2	2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
C	2	2.7.8	When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison testing implemented. The results are recorded and this record is maintained.

K	2, 11		2.7.9 New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2		2.7.10 The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2		2.7.11 Membrane filters which are beyond their expiration date are not used.
O	11		2.7.12 Forceps tips are clean.
O	11		2.7.13 Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.
K	11		2.7.14 Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11		2.7.15 If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11		2.7.16 Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11		2.7.17 Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26		2.7.18 A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11		2.7.19 The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2		2.7.20 Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
2.8 Media Preparation and Storage – MF using mTEC Agar			
K	11		2.8.1 Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
C	11		2.8.2 The phosphate buffered saline is properly sterilized.
K	23		2.8.3 A sufficient amount of medium (4-5 mL) is used in each plate.
O	11		2.8.4 Refrigerated prepared plates are stored for no more than 2 weeks in sealed plastic bags or containers to minimize evaporation.
2.9 Sample Analyses - MF using mTEC Agar			
C	24		2.9.1 mTEC agar is used.
C	2		2.9.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control
C	23, 35		2.9.3 The sample is shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration
C	23		2.9.4 The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25		2.9.5 Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23		2.9.6 Sample volumes are filtered under vacuum.
K	26		2.9.7 The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26		2.9.8 The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.
C	23		2.9.9 The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11		2.9.10 Blanks are run at the beginning of filtration, after every 10th aliquot and at the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).

C	2, 11		2.9.11 Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24		2.9.12 Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at 35 + 0.5°C for 2 hours of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5°C for 24 ± 2 hours.
C	11, 23, 24		2.9.13 After 2 hours of resuscitation at 35°C, the watertight, tightly sealed containers are transferred to a circulating waterbath at 44.5 + 0.2°C, submerged completely and incubated for 22-24 hours.
			2.10 Computation of Results - MF using mTEC Agar
C	23		2.10.1 All yellow, yellow-green or yellow-brown colonies are counted.
C	23		2.10.2 Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as >80 x 100/the volume of sample filtered.
C	2, 11, 23		2.10.3 When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11		2.10.4 The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11		2.10.5 Results are reported as CFU/100 mL of sample.
			2.11 Bacteriological Analysis of UV Treated Process Water Samples by Membrane Filtration (MF) using mEndo Agar LES – Materials and Equipment
C	9, 11, 21		2.11.1 Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches are used.
C	2		2.11.2 The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
C	11, 19, 21		2.11.3 Membrane filters are made from cellulose ester material, white, grid marked, 47 mm diameter with a pore size of 0.45µm and certified by the manufacturer for total coliform analysis.
C	2		2.11.4 Membrane filter lot numbers, dates of receipt and expiration dates are recorded and the records maintained.
C	2		2.11.5 If previous lots of agar or membrane filters are not available for comparability testing, an appropriate method for determining lot suitability has been developed and comparison testing implemented when the following has occurred: a. initiating monitoring by mEndo Agar LES; b. changing agar manufacturers; or c. changing brands of membrane filters used. The results are recorded and the records are maintained.
K	2, 11		2.11.6 Recovery of total coliforms from new lots of membrane filters and mEndo Agar LES is compared against the recovery from the previously acceptable lot.
C	2		2.11.7 The sterility of each lot and autoclave batch of membrane filters is verified before use.
K	2		2.11.8 Expired membrane filters are not used.
K	9, 11, 19, 21		2.11.9 Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.

K	11		2.11.10 If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
C	9, 11		2.11.11 Membrane filter assemblies are autoclave sterilized for 15 minutes at 121+/- 2°C prior to the start of a filtration run. A new run occurs when there is a break of 30 minutes or more between the previous filtration run.
O	11, 19, 26, 36		2.11.12 A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11		2.11.13 The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and the records are maintained.
K	2		2.11.14 Maintenance of the UV sterilization unit is performed as needed. Maintenance is documented and the records maintained.
O	9, 11		2.11.15 Forceps tips are clean and smooth without pitting or corrugations.
			2.12 Media Preparation and Storage
C	9, 11, 19, 21, 36		2.12.1 mEndo Agar LES is used.
K	11, 21, 36		2.12.2 mEndo Agar LES is prepared aseptically using pre-sterilized glassware, sterile reagent water and pre-sterilized stir bar.
K	9, 11, 36		2.12.3 mEndo Agar LES is prepared using 95% alcohol that is not denatured.
C	9, 11, 36		2.12.4 mEndo Agar LES is brought to near boiling; removed from the heat and tempered at 44-50°C before dispensing.
C	9, 11, 36		2.12.5 mEndo Agar LES is never autoclaved.
K	9, 11, 36		2.12.6 A sufficient amount of medium (4-5 mL) is aseptically aliquoted to each culture plate.
O	9, 11, 36		2.12.7 Prepared plates of mEndo Agar LES are stored at 4°C in the dark for no more than two (2) weeks in sealed plastic bags or containers to minimize evaporation.
C	2		2.12.8 Appropriate, properly diluted positive and negative productivity controls for mEndo Agar LES medium are used. Results are recorded and the record maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11, 21, 36		2.12.9 Sterile phosphate buffered water or sterile phosphate buffered saline is used as a sample blank, filter funnel rinse and process and productivity control diluent for UV treated process water samples.
C	11		2.12.10 The phosphate buffered water/saline is properly sterilized and the sterility is tested before being placed in service. Results are recorded and records maintained
			2.13 Sample Analysis
C	9, 11, 36		2.13.1 The sample is shaken vigorously (25 times in a 12-inch arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration.
C	11, 21, 36		2.13.2 The membrane filter is placed grid side up within the sterile filter apparatus.
C	11, 26, 36		2.13.3 A 100 mL quantity of sample is filtered under vacuum.
K	26		2.13.4 The pressure of the vacuum pump does not exceed 15 psi.
C	9, 11, 26, 36		2.13.5 The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered water/saline as appropriate after filtration.
C	9, 11, 36		2.13.6 The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mEndo Agar LES so that no bubbles form between the filter and the agar.

K	9, 11, 36		2.13.7 Forceps are dipped in alcohol and flame sterilized between sample filters.
C	11, 36		2.13.8 Blanks are run at the beginning and at the end of the filtration run to check the sterility of the testing system (phosphate buffered water/saline, filter funnels, forceps, membrane filters, media and culture plates).
C	2, 36		2.13.9 An appropriate properly diluted positive process control culture accompanies the sample throughout incubation. Results are recorded and the records are maintained. Positive process control _____
C	9, 11, 36		2.13.10 Inoculated plates are incubated inverted at 35+/- 0.5°C for 22 to 24 hours.
K	2, 9, 11		2.13.11 An appropriate level of humidity is maintained in the incubator to prevent the plates from drying out.
2.14 Computation of Results			
K	9, 11		2.14.1 Colonies are counted with the aid of magnification.
C	9, 11, 19, 21, 23		2.14.2 All metallic sheen colonies are counted as total coliforms.
C	9, 11, 21, 36		2.14.3 Results are reported as total coliforms/100mL.
C	11, 20, 36		2.14.4 When no colonies are observed, results are reported as <1.0 coliform/100mL (nondetectable)
PART III - SHELLFISH SAMPLES			
3.1 Collection and Transportation of Samples			
C	9		3.1.1 A representative sample of shellstock is collected.
K	9		3.1.2 Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9		3.1.3 Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9		3.1.4 Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1		3.1.5 Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
3.2 Preparation of Shellfish for Examination			
K	2,11		3.2.1 Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2		3.2.2 Blades of shucking knives are not corroded.
O	9		3.2.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2		3.2.4 The faucet used for rinsing the shellstock does not contain an aerator.
K	9		3.2.5 Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9		3.2.6 Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	1, 9		3.2.7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9		3.2.8 Shellstock are not shucked directly through the hinge.
C	9		3.2.9 Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9		3.2.10 At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.

K	9	3.2.11	A representative sample of at least 12 shellfish is used for the analysis.
K	2	3.2.12	The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
Ø	9	3.2.13	Sterile phosphate buffered dilution water is used as the sample diluent.
C	9	3.2.14 13	Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9	3.2.15 14	APHA Recommended Procedures for the Examination of Sea Water And Shellfish , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA			
C	9	3.3.1	Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. (Circle the medium used.)
C	2	3.3.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9	3.3.3	Immediately (within 2 minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9	3.3.4	No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
C	9	3.3.5	Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6	3.3.6	In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	3.3.7	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9	3.3.8	Inoculated media are incubated at $35 \pm 0.5^\circ\text{C}$.
K	10	3.3.9	Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA			
C	9	3.4.1	EC medium is used as the confirmatory medium.
C	2	3.4.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11	3.4.3	Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. (Circle the method of transfer.)
C	9	3.4.4	EC tubes are incubated in a circulating waterbath at $44.5 \pm 0.2^\circ\text{C}$
K	9	3.4.5	EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9	3.4.6	The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.
3.5 Computation of Results for MPN Analyses			
K	9	3.5.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.

K	7		3.5.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9		3.5.3	Results are reported as MPN/100 grams of sample.
			3.6 Standard Plate Count Method	
O	20		3.6.1	A standard plate count (SPC) analysis maybe performed in conjunction with the analysis for fecal coliform organisms.
K	9		3.6.2	In the standard plate count procedure at least four plates are used, duplicates of two dilutions. One of the dilutions should produce colonies of 30 to 300 per plate.
K	2		3.6.3	Fifteen to 20 mL of tempered sterile plate count agar is used per plate.
C	9		3.6.4	Agar tempering bath maintains the agar at 44-46°C.
C	9		3.6.5	An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9		3.6.6	Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in 7 seconds) before plating.
C	9		3.6.7	Not more than 1 mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11		3.6.8	Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9,21		3.6.9	Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four high.
K	9		3.6.10	Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1		3.6.11	A hand tally or its equivalent is used for accuracy in counting.
			3.7 Computation of Results -SPC	
K	9		3.7.1	Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19		3.7.2	Colony counts are reported as CFU/g of sample.
			3.8 Bacteriological Analysis of Shellfish Using the ETCP	
C	2,3		3.8.1	Prepared modified MacConkey agar is used on the day that it is made.
K	3		3.8.2	Double strength modified MacConkey agar is used.
C	3		3.8.3	Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3		3.8.4	Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50°C until used.
K	2, 3		3.8.5	Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3		3.8.6	The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9		3.8.7	The sample homogenate is cultured within 2 minutes of blending.
C	2,3		3.8.8	Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.
K	3		3.8.9	Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2,3, 22		3.8.10	The container is gently swirled or slowly inverted once to mix the contents, which are subsequently distributed uniformly over six plates.
C	1		3.8.11	Media and diluent sterility are determined with each use. Results are recorded and the records maintained.

C	1		3.8.12 Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture _____ Negative control culture _____
C	3, 13		3.8.13 When solidified, the plates are placed inverted into an air incubator at 45.5 ± 0.5°C for 18 to 30 hours of incubation.
C	2		3.8.14 Plates are stacked no more than three high in the incubator.
C	2		3.8.15 Appropriately diluted pour plated process control cultures accompany each set of samples throughout incubation. The results are recorded and the records maintained. Positive process control _____ Negative process control _____
			3.9 Computation of Results - ETCP
K	11		3.9.1 Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1		3.9.2 A hand tally or its equivalent is used to aid in counting.
C	3, 6		3.9.3 All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3		3.9.4 Results are reported as CFU/100 grams of sample.
			Bacteriological Examination of Soft-shelled Clams and American Oysters for Male Specific Coliphage (MSC)
			3.10 MSC Equipment and Supplies
K	30		3.10.1 Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold 100 – 125 mL.
C	27, 28		3.10.2 The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	9		3.10.3 The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
C	27, 28		3.10.4 Sterile 0.22µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1		3.10.5 The sterility of each lot of pre-sterilized syringes and syringe filters is determined. Results are recorded and records maintained.
K	1		3.10.6 The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28		3.10.7 The balance used provides a sensitivity of at least mg (0.01g.).
C	27, 28		3.10.8 The temperature of the incubator used is maintained at 36 ± 1°C.
C	28		3.10.9 Sterile disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
			3.11 MSC Media Preparation
K	28		3.11.1 Media preparation and sterilization is according to the validated method.
K	27, 28		3.11.2 Bottom agar, double strength soft agar and growth broth are prepared from their individual components.
K	27, 28		3.11.3 Soft agar is prepared double strength in volumes of 2.5 mL.
C	27, 28		3.11.4 The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
O	27, 28		3.11.5 Storage of the bottom agar under refrigeration does not exceed 1 month.
K	27, 28		3.11.6 Unsterilized soft agar is stored at -20 °C -15C for up to 3 months.
K	27, 28		3.11.7 The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.
K	27, 28		3.11.8 Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	27, 28		3.11.9 Bottom agar plates are allowed to reach room temperature before use.

3.12 Preparation of the Soft-Shelled Clams and American Oysters for MSC Analysis			
K	2,11		3.12.1 Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2		3.12.2 The blades of shucking knives are not corroded.
O	9		3.12.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2		3.12.4 The faucet used for rinsing the shellfish does not contain an aerator.
K	9		3.12.5 The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9		3.12.6 The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9		3.12.7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9		3.12.8 Shellfish are not shucked through the hinge.
C	9		3.12.9 The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9		3.12.10 A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19		3.12.11 The sample is weighed to the nearest 0.1 gram.
3.13 MSC Sample Analysis			
C	28		3.13.1 <i>E. coli</i> Famp ATCC 700891 is the bacterial host strain used in this procedure.
K	27, 28		3.13.2 Host cell growth broth is tempered at $36 \pm 1^\circ\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	27, 28		3.13.3 Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^\circ\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	27, 28		3.13.4 After inoculation, the host cell growth broth culture is not shaken.
C	28		3.13.5 A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.
C	28		3.13.6 The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28		3.13.7 The elution mixture is homogenized at high speed for 180 seconds.
C	28		3.13.8 Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
C	28		3.13.9 The homogenized elution mixture is centrifuged for 15 minutes at $9000 \times g$ at 4°C.
C	27, 28		3.13.10 The supernatant is pipetted off, weighed and the weight recorded.
C	27, 28		3.13.11 The supernatant is allowed to warm to room temperature about 20 to 30 minutes.
K	27, 28		3.13.12 The autoclaved soft agar is tempered and held at $51 \pm 1^\circ\text{C}$ throughout the period of sample analysis.
K	27, 28		3.13.13 Two hundred microliters (0.2 mL) of log phase host strain <i>E. coli</i> is added to the tempering soft agar immediately prior to adding the sample supernatant.
K	27, 28		3.13.14 The sample supernatant is shaken or vortexed before being added to the tempering soft agar.
C	27, 28		3.13.15 2.5 mL of sample supernatant is added to each tube of tempering soft agar.
C	27, 28		3.13.16 The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28		3.13.17 The soft agar/sample supernatant/host cell mixture is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.
C	28		3.13.18 Ten (10) plates are used, 2.5 mL per plate for a total of 25 mL of supernatant analyzed per sample.

K	27, 28		3.13.19 Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are recorded and records maintained. Positive control _____
K	27, 28		3.13.20 Growth broth is used as the negative control or blank.
K	27, 28		3.13.21 Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2		3.13.22 A negative control plate is plated at the beginning and end of each set of samples analyzed.
K	27, 28		3.13.23 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28		3.13.24 All plates are incubated at 36 ± 1°C for 18 ± 2 hours.
3.14 Computation of Results - MSC			
C	27		3.14.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 32		3.14.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten plates, the count is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/ 100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as > 20,000 PFU/100 grams.
K	28		3.14.3 The formula used for determining the density of MSC in PFU/100 grams is: (0.364) (N) (Ws), where N = total number of plaques counted on all 10 plates and Ws = weight of the supernatant used.
O	9		3.14.4 The MSC count is rounded off conventionally to give a whole number.

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PART I – Quality Management: Laboratory Operations and Responsibilities for National Shellfish Sanitation Program Laboratory Quality Systems		
ITEM		
Conformance Comments	Ref	
		1.1 Components of the Laboratory Quality System
	1,3,6,9	1.1.1 The laboratory has an overall Quality System supported by quality management structure, quality assurance processes and quality control functions.
	1,3,6,9	1.1.2 Management and technical structure exist to support the Quality System.
	1,3,6,9	1.1.3 Quality documentation is required by the laboratory. These include a Quality Assurance (QA) Manual (or otherwise named) and Standard Operating Procedures (SOPs) to support the quality assurance process of the laboratory.
	1, 9	1.1.4 The documents used to implement the quality assurance process and records used to verify quality control (QC) function of the laboratory are reviewed and controlled.
	9	1.1.5 An established process of Quality System assessment and technical proficiency are documented with results retained until the next review.
	9	1.1.6 Resolution, management review and prevention of nonconformities are a documented component of the Quality System.
		1.2 Laboratory Management Structure and Quality Systems
	1,3,6,9	1.2.1 The laboratory's structure is clearly organized with supervisory chain delineated.
	9	1.2.2 The laboratory has ensured that its management and personnel are free from any undue internal and external commercial, financial and other pressures and influences that may adversely affect the quality of their work.
	9	1.2.3 The laboratory has documentation of dedicated <u>designated quality personnel and/or a designated quality manager</u> with the authority and resources required to carry out their duties, including implementing and maintaining the Quality System, of the laboratory. <u>ensure adherence to the Quality System, initiate actions to prevent or minimize departures from the Quality System, and monitor all aspects of the Quality System to ensure defensibility. This person has unrestricted access to FDA Shellfish Laboratory Evaluation Officers (LEOs) and the highest levels of the laboratory's management. In the case of a single person laboratory, FDA LEOs will assist with developing a monitoring plan.</u>
	1, 9	1.2.4 The laboratory's designated quality personnel ensure adherence to the quality system, including SOPs and QC. These staff have clear documented authority to initiate actions to prevent or minimize departures from quality system and monitor the corrective action process.
	9	1.2.5 The laboratory has documentation of a designated quality system manager, responsible for monitoring all aspects of the quality system to assure defensibility. This person shall have unrestricted access to FDA Shellfish Laboratory Evaluation Officers (LEOs) and the highest levels of the laboratories management. In the case of a single person laboratory, FDA LEOs will assist with developing a monitoring plan.

	1, 9	1.2.64 A documented system is in place to ensure that appropriate review of and communication regarding the elements of the quality system are established among the laboratory staff and laboratory management.
		1.3 Laboratory Personnel and Roles in a Quality System
	1,3, 9	1.3.1 The roles and responsibilities of all personnel are defined in the QA manual, read by all staff and the acknowledgments of these responsibilities are retained.
	9	1.3.2 The laboratory policy and the training procedures for personnel are documented and relevant to the scope of the current activities in the laboratory. If the laboratory intends to add methods to their scope, training SOPs must also be added with successful completion by the analyst(s) that will perform the method(s). In the case of a single person laboratory, method proficiency verification must be retained during the life of the methods use in the laboratory.
	9	1.3.3 The laboratory shall maintains a personnel file/ record of any relevant authorization(s), qualifications, trainings, and/or proficiencies for each analyst. This information shall be is available upon request as verification of staff training and shall be is retained for all staff until two years after they are no longer employed by the laboratory.
	1, 3, 9	1.3.4 The laboratory has documented that all personnel involved in testing have read and understand the applicable SOPs and associated quality documentation and implement the policies and procedures required for the performance of their technical function.
PART II – Quality Assurance: The Process of Documenting and Maintaining a Quality System		
		2.1 Quality Assurance Process: QA Manual, SOPs and Document Control
	1, 9	2.1.1 The QA manual shall includes or make references to all laboratory SOPs and any supporting procedures, including technical procedures.
	1, 9	2.1.2 SOPs are controlled documents and include detailed, written instructions to achieve uniformity of test methods and quality control procedures, such that items that might affect the quality or defensibility of the outcome are mitigated.
	1, 9	2.1.3 SOPs and the QA Manual are controlled documents, such that specific individuals are designated within the laboratory with editorial control. These individuals are identified in the QA Manual.
	1, 9	2.1.4 Each time an SOP or the QA manual has changed, the new version will be marked as such and will be distributed to the laboratory with older versions removed from circulation.
	1, 9	2.1.5 Staff training requirements are documented in the QA manual and the training procedure is included.

2.2 Quality Manual Items		
1, 9	2.2.1	Quality Assurance Manual contains: <ol style="list-style-type: none"> 1. Table of Contents; 2. Organizational chart; 3. A description of the Quality System and procedure for implementation and maintenance; 4. Policy and procedure for resource management (human resources, competence and training, work environment and safety), description of responsibilities; 5. Policy and procedures for rejection criteria; 6. Policy and procedures for calibration of equipment and Equipment file items such as maintenance; 7. Policy and procedure for traceability and required documentation, 8. Policy and procedure for internal audits; 9. Policy and Procedure for data analysis and control of nonconforming work; and 10. Policy for corrective actions (CAs) and preventative actions (PAs).
1,3,6,9	2.2.2	The organizational chart clearly depicts laboratory structure with quality and technical personnel listed.
1, 9	2.2.3	The policy for human resources provisions includes hiring and assignment of staff, competence and responsibilities for positions, and a procedure of training for each technical competence, including proficiencies required.
1, 3, 4, 6, 9	2.2.4	Policies for work environment and safety protocols, analytical methods, and quality control performed for the National Shellfish Sanitation Program (NSSP) are included or referenced in the QA Manual and shall be provided upon request.
1, 9	2.2.5	A policy regarding appropriate equipment file maintenance and retention (e.g., calibration records, maintenance documentation, manuals of operation) is included in the QA Manual.
1, 9	2.2.6	The SOP for calibration and maintenance of equipment is kept or referenced in the QA Manual and shall be provided upon request.
1, 9	2.2.7	The SOP for traceability of analytical results is included or referenced in the QA Manual and shall be provided upon request. This traceability procedure includes a documented procedure for the unique identification of samples and the process for chain of custody verification.
1, 9	2.2.8	The QA Manual has a policy and a procedure for internal quality audits. These audits are planned and scheduled annually or as needed. The policy states auditors do not audit their own work. In the case of a single person laboratory, FDA LEOs will assist with an audit plan.
1, 9	2.2.9	The QA Manual contains a policy for data analysis to require that all analyses performed have been carried out correctly, documented, controls were used accurately and the results meet specified requirements.

	1, 9	2.2.10 The QA Manual contains a procedure for the control of nonconforming work in the case of: <u>a.</u> identification, documentation, evaluation, segregation (where practical), disposition of nonconforming sample/analyte/result and customer notification; <u>b.</u> assigning responsibility for the review and the authority for disposition of nonconforming sample/analyte/result; <u>c.</u> a nonconforming result correction and the re-verification/calibration of the affected equipment after the correction to demonstrate conformity (if necessary); and <u>d.</u> handling a nonconforming result when it is detected, after delivery to the customer.
	1, 9	2.2.11 The QA manual contains a procedure for preventative actions in which laboratory staff identify potential nonconformities in audit results, quality records, or customer complaints through a review process. Steps are then determined to identify preventive actions to implement. The necessary changes are made to SOPs and this exercise is recorded, and records maintained.
	1, 3, 6, 9	2.2.12 The QA manual has a policy and a procedure for developing corrective action(s) to eliminate the cause of identified nonconformities in order to prevent recurrence. Corrective actions describe the nonconformities, define the process for evaluating the need for actions to ensure that nonconformities do not recur (root cause analysis), explain the process to implement the corrective action(s) needed, and the resultant outcome. There is also a procedure to monitor progress of any ongoing corrective actions and the resolution.
	1, 3, 4, 6, 9	2.2.13 The QA Manual contains a policy stating laboratory management shall ensure and document the competence of staff independently operating equipment resulting in a documented measurement, analysis result, quality control value/result, determination of data value for sample result, and review/closure of corrective action for efficacy.
	1, 9	2.2.14 The policy for sample rejection criteria includes what the laboratory will accept and reject based on NSSP requirements and chain of custody.
	1, 3, 4, 6, 9	2.2.15 The laboratory shall have <u>has</u> sample acceptance procedures that include safe handling, transport, and storage to prevent contamination or deterioration and to protect the sample integrity. These procedures are provided to customers.
	1, 3, 4, 6, 9	2.2.16 The laboratory has procedures for handling nonconforming samples and who will be contacted in the case of sample rejection.
PART III- Quality Control: Documentation for Quality System Defensibility		
3.1 Documentation		
	1, 9	3.1.1 The laboratory investigates proficiency testing (PT) programs for areas of continual improvement and actively addresses problematic results through the prescribed corrective action process.
	1, 9, 10	3.1.2 The laboratory personnel performing sampling and testing sample <u>analyses</u> participate in PT programs and exercises when available. If no PT exists, participation in interlaboratory comparisons is considered.

1, 3, 6, 9, 10	3.1.3	Corrections to quality control records, bench sheets and reports follow the requirements below: <u>a.</u> A single line is drawn through the incorrect information; <u>b.</u> The correct information is written next to the incorrect information; <u>c.</u> The person responsible for the correction initialed the information; <u>d.</u> If not obvious, the reason for correction has been included; and <u>e.</u> If corrections are necessary in an electronic document, old information must be retained in some form, the person making the change must be identified, the date of the change noted, and the reason for the change noted.
1, 3, 6, 9, 10	3.1.4	All records, required to be retained for two years (or length of time as dictated by State law), shall be <u>are</u> legible and shall be stored in such a way that they are readily retrievable to prevent damage or loss.
1	3.1.5	All records and documents must be written in indelible ink.
		3.2 Method Performance Validation <u>Verification</u>
1, 3, 6, 9	3.2.1	The laboratory will internally validate <u>verify</u> new methods to confirm with objective evidence that the intended protocols are demonstrated and outcomes are fulfilled.
1, 9	3.2.2	Methodologies do not deviate from the validated/ <u>verified</u> method and the laboratory's internal validation shall <u>verification</u> remains on file in the laboratory.
1, 3, 6, 9, 10	3.2.3	The laboratory shall <u>reports</u> the method chosen in writing to the customer.
1, 4, 9	3.2.4	Methodologies and protocols are selected <u>and samples are processed</u> based on NSSP requirements, and samples are processed as per the citation in the current Model Ordinance.
		3.3 Environmental Conditions
1, 3, 4, 5, 6, 9, 10	3.3.1	Laboratory facilities <u>support accurate test performance for analysis</u> , including lighting and environmental conditions such as temperature and humidity, shall support accurate performance of the tests .
1, 3, 4, 5, 6, 9, 10	3.3.2	The laboratory shall <u>monitors</u> , controls, and records environmental conditions as required by the relevant specifications, methods, and procedures, or where they influence the outcome of results (e.g., biological sterility, dust, humidity, electrical supply, temperature, vibration).
1, 3, 4, 6, 9, 10	3.3.3	Laboratory personnel shall <u>stop</u> testing when the environmental conditions jeopardize the results of analyses.
1, 3, 4, 6, 9, 10	3.3.4	Personnel shall <u>ensure</u> good housekeeping in the laboratory.
		3.4 Equipment
1, 3, 4, 6, 9, 10	3.4.1	The laboratory shall have <u>has</u> instructions and/ or SOPs on the use and operation of all relevant equipment, and on the handling and preparation of items for testing, where the absence of such could jeopardize the outcome of analysis or influence results.
1, 9, 10	3.4.2	All equipment in the laboratory is labelled with the manufacturer's name, identification number, and serial number or other unique identification that is traceable.
1, 9, 10	3.4.3	Equipment files contain reports and certificates of all calibrations, the due date of next calibration, dates and results of any maintenance, adjustments, damage, malfunction, and modification or repair to the equipment.

	1, 2, 9, 10	3.4.4 If equipment (e.g., thermometer, balance) was sent out of the laboratory for service, performance has been verified prior to use again in the laboratory.
		3.5 Temperature Measuring Devices
	1, 2, 8, 9, 10	3.5.1 Unique identifier, <u>verification or calibration</u> ice point date (if applicable) and any correction factor is recorded on <u>each</u> in use temperature measuring device (TMD).
	1, 2, 8, 9, 10	3.5.2 TMDs are calibrated <u>and verified</u> as per the NSSP requirements and ice points/steam points are performed annually on Standards thermometers.
	1, 8	3.5.3 TMDs calibration certificates are retained for three consecutive calibration cycles (<u>as applicable</u>).
	1, 8, 9, 10	3.5.4 Where calibrations give rise to a set of correction factors, the laboratory shall have <u>has</u> procedures to ensure <u>these correction factors are appropriately applied to equipment and</u> records are retained until the next <u>verification check</u> is performed.
	1, 8, 9, 10	3.5.5 <u>Accuracy, Range,</u> and graduations of all TMDs are appropriate for the designated use. Dial thermometers are not used in the laboratory.
	8, 9, 10	3.5.6 For electronic TMDs, probe/sensor is uniquely labeled and placement within unit being monitored follows manufacturer's instructions <u>are followed</u> to ensure accurate readings, as devices vary.
	1, 8, 9, 10	3.5.7 Temperature Monitoring Systems (wired/wireless) must record temperature <u>reading</u> from each sensor/probe in the piece of equipment being monitored at the same or greater frequency <u>required by the NSSP.</u> and accuracy as stipulated for mercury in-glass thermometers, as per manufacturer specifications.
		3.6 Disposables and Pipettors
	1, 3, 4, 6, 9, 10	3.6.1 Pipettors, accuracy checked, fixed volume or electronic are calibrated according to NSSP requirements.
	1, 3, 10	3.6.2 Pipettors are etched <u>or imprinted</u> with <u>unique</u> identification (imprinted serial numbers acceptable) and tagged with last date of accuracy <u>check</u> verification.
	1, 3, 4, 6, 9, 10	3.6.3 Appropriate pipettor tips are used and sterility checks are performed on an appropriate quantity.
	1, 3, 4, 6, 9, 10	3.6.4 Sterility checks on disposables are performed according to a cited QC practice, within a designated SOP; (e.g., laboratory may cite and implement a recognized standard of sterility testing, they may test 10% of a "lot" or any 3 in a box).
		3.7 Test Record/Bench Sheet Requirements
	1, 3, 4, 6, 9, 10	3.7.1 Test records/bench sheets shall contain information to facilitate repeatability under conditions as close as possible to the original including QC information (or reference) for media and supplies used.
	1, 9, 10	3.7.2 Test records/bench sheets must show date, time and temperature of samples at the start of analysis and contain the name or initials of the analyst performing the test for each group of samples.
	1, 4, 9, 10	3.7.3 Test records/bench sheets must include sterility controls or a reference to the document containing sterility controls for disposables and dilution buffer.
	1, 4, 9, 10	3.7.4 Test records/bench sheets must include media productivity (positive and negative) controls or a reference to the document containing media productivity controls.

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PART 1 - QUALITY ASSURANCE		
CODE	REF.	ITEM
K	8, 11	1.1 Quality Assurance (QA) Plan
		1.1.1 Written Plan (Check those items which apply.)
		a. Organization of the laboratory.
		b. Staff training requirements.
		c. Standard operating procedures.
		d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		e. Laboratory safety.
		f. Internal performance assessment.
		g. External performance assessment.
C	8	1.1.2 QA Plan Implemented.
K	11	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s) _____
		1.2 Educational/Experience Requirements
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three months of experience in laboratory science
		1.3 Work Area
O	8,11	1.3.1 Adequate for workload and storage.
K	11	1.3.2 Clean, well-lighted.
K	11	1.3.3 Adequate temperature control.
O	11	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
		1.4 Laboratory Equipment
O	9	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	1.4.4 pH meter is calibrated daily or with each use. Results are recorded and records maintained.
K	11	1.4.5 A minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	8,15	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt procedure or through determination of the slope. (<i>Circle the method used.</i>)

K	9	1.4.7	Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13	1.4.8	Balance calibrations are verified checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11	1.4.9	Refrigerator temperature(s) are monitored at least once daily on workdays Results are recorded and records maintained.
K	1	1.4.10	Refrigerator temperature is maintained at 0 to 4°C.
C	9	1.4.11	The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11	1.4.12	Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9	1.4.13	Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11	1.4.14	Temperature of the waterbath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9	1.4.15	The thermometers used in the waterbath are graduated in at least 0.1°C increments.
C	13	1.4.16	The waterbath has adequate capacity for workload.
K	9	1.4.17	The level of water in the waterbath covers the level of liquid in the incubating tubes.
K	8, 11	1.4.18	Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4	1.4.19	All working thermometers are appropriately immersed.
C	29	1.4.20	Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	11	1.4.21	A standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9	1.4.22	Standards thermometers are verified checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination_____
C	29	1.4.23	Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers <u>with an accuracy and tolerance appropriate for the application</u> with an accuracy of ≤±0.05°C are used as the laboratory standards thermometer. (Circle the thermometer type used.)
K	13	1.4.24	The accuracy of working thermometers is checked annually against the standards thermometer either at the temperatures at which they are used or by ice point determination. Results are recorded and records maintained.
O	11	1.4.25	Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing			
O	9	1.5.1	Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding materials.
K	9	1.5.2	Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.
K	9	1.5.3	Sample containers are made of glass or some other inert material.
O	9	1.5.4	Dilution bottles and tubes are made of borosilicate glass or plastic and closed with rubber stoppers, caps or screw caps with nontoxic liners.

K	9	1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9	1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1mL used to deliver 0.1 mL aliquots.
K	9	1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9	1.5.8	In washing reusable pipettes, a succession of at least three fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2	1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11	1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination			
K	9	1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8	1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	30, 33, 34	1.6.3	The autoclave provides sterilization conditions suitable to the load contents. Sterilization temperature range may be 119°C - 124°C as determined by the lab's equipment Quality Assurance Verification Testing and recommended practices from the media manufacturer. Sterilization is determined for each load using a working maximum registering thermometer or an appropriate working temperature monitoring device.
K	11	1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a <u>device traceable primary standard traceable</u> to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16	1.6.5	The autoclave standards thermometer is <u>recalibrated</u> checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory; or, is <u>verified</u> checked in-house at the steam point (100°C) if it <u>was</u> has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination _____
K	1	1.6.6	Working autoclave thermometers are <u>verified</u> checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11	1.6.7	Spore strips/suspensions <u>with a kill time</u> appropriate for use in an autoclave <u>liquid</u> media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11	1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13	1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11	1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180°C.
K	9	1.6.11	A thermometer capable of determining temperatures accurately in the range of 160 to 180°C is used to monitor the operation of the hot-air sterilizing oven
K	13	1.6.12	Records of temperatures and exposure times are maintained for the operation of the hot-air sterilizing oven during use.

K	11		1.6.13 Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Records are maintained.
K	11		1.6.14 Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C.
C	1		1.6.15 The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1		1.6.16 The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9		1.6.17 Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9		1.6.18 Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2 hours.
C	2		1.6.19 The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2		1.6.20 The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18		1.6.21 Hardwood applicator transfer sticks or <u>reusable loops</u> are properly sterilized. <u>Alternatively, presterilized loops are used for transfers.</u> Method of sterilization _____
C	2		1.6.22 The sterility of the hardwood applicator transfer sticks/<u>presterilized loops</u> is checked routinely. Results are recorded and the records maintained.
O	13		1.6.23 Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation			
K	3, 5		1.7.1 Media is commercially dehydrated except in the case of medium A-1 which must be prepared from the individual components and modified MacConkey agar which may be prepared from its components.
K	11		1.7.2 Media is prepared <u>and sterilized</u> according to manufacturer's instructions.
O	11		1.7.3 Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11		1.7.4 Dehydrated media are labeled with date of receipt and date opened.
C	12		1.7.5 Caked or expired media or media components are discarded.
C	11		1.7.6 Reagent water is distilled or deionized (circle appropriate choice), tested monthly and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) or is less than 2.0 µSiemens/cm conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11		1.7.7 Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level (<0.1 mg/L). Results are recorded and the records maintained. Specify method of determination_____.
K	11		1.7.8 Reagent water contains <100 CFU/mL as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11		1.7.9 Media prepared from commercial dehydrated components are sterilized according to the manufacturer's instructions.
K	9		1.7.9+0 The volume and concentration of media in the tube are suitable for the amount of sample inoculated.
C	11		1.7.10+ Total time of exposure of sugar broths to autoclave temperatures does not exceed 45 minutes.

C	1	1.7.11² Media sterility is determined for each load sterilized. Results are recorded and the records maintained.
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C	1		1.7.13³ Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is made from its individual components.
O	9		1.7.1 3 ⁴ Sterile phosphate buffered dilution water is used as the sample diluent.
K	11		1.7.1 4 ⁵ The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media			
K	9		1.8.1 Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11		1.8.2 Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13		1.8.3 Stored media are labeled with the storage expiration date or the sterilization date.
K	9		1.8.4 Storage of prepared culture media at room temperature does not exceed 7 days.
K	2		1.8.5 Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed 1 month.
K	11		1.8.6 Storage under refrigeration of prepared culture media with screw-cap closures does not exceed 3 months.
K	17		1.8.7 All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES			
2.1 Collection and Transportation of Samples			
C	11		2.1.1 Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1		2.1.2 Samples are identified with collector's name, harvest area, sampling station, time and date of collection.
C	9		2.1.3 Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1		2.1.4 A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9		2.1.5 Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.
2.2 Bacteriological Examination of Seawater by the APHA MPN			
C	9		2.2.1 Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
C	2		2.2.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _
C	9, 35		2.2.3 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9		2.2.4 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6		2.2.5 In a single dilution series not less than 12 tubes are used (for depuration at least 5 tubes are used).

C	6		2.2.6 In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
K	9		2.2.7 Inoculated tubes are incubated in air at 35 ± 0.5°C.
C	2		2.2.8 Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
K	9		2.2.9 Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
2.3 Confirmed Test for Seawater by APHA MPN			
C	9		2.3.1 Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9		2.3.2 EC medium is used as the confirmatory medium for fecal coliforms.
C	2		2.3.3 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11		2.3.4 Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. (Circle the method of transfer.)
C	9		2.3.5 BGB tubes are incubated at 35 ± 0.5°C.
K	9		2.3.6 BGB tubes are read after 48 ± 3 hours of incubation.
C	9		2.3.7 EC tubes are incubated in a circulating waterbath maintained at 44.5 ± 0.2°C.
C	9		2.3.8 EC tubes are read after 24 ± 2 hours of incubation.
C	9		2.3.9 The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.4 Computation of Results – APHA MPN			
K	9		2.4.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7		2.4.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9		2.4.3 Results are reported as MPN/100 mL of sample.
2.5 Bacteriological Examination of Seawater by the MA-1 Method			
C	5		2.5.1 A-1 medium complete is used in the analysis.
C	2, 31		2.5.2 A-1 medium without salicin is used in the analysis. Comparability testing supports use of A-1 medium without salicin. Study records are available.
C	5		2.5.3 A-1 medium sterilized for 10 minutes at 121°C.
C	2		2.5.4 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____

C	9, 35	2.5.5	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9	2.5.6	In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	2.5.7	In a single dilution series at least 12 tubes are used.
C	6	2.5.8	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	2.5.9	Appropriately diluted process control cultures accompany the samples <i>throughout both resuscitation and waterbath incubation</i> Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	2.5.10	Inoculated tubes are placed in an air incubator at $35 \pm 0.5^\circ\text{C}$ for 3 ± 0.5 hours of resuscitation.
C	5	2.5.11	After 3 ± 0.5 hours resuscitation at 35°C , inoculated tubes are incubated at $44.5 \pm 0.2^\circ\text{C}$ in a circulating waterbath for the remainder of the 24 ± 2 hours.
C	5	2.5.12	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.6 Computation of Results – APHA MPN			
K	9	2.6.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	2.6.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.6.3	Results are reported as MPN/100 mL of sample.
2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment			
C	23, 24	2.7.1	When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at $44.5 \pm 0.5^\circ\text{C}$ under any loading capacity.
C	23	2.7.2	When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	2.7.3	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	2.7.4	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	2.7.5	Colonies are counted with the aid of magnification.
C	11, 23	2.7.6	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of 0.45 μm and certified by the manufacturer for fecal coliform analyses.
C	2	2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
C	2	2.7.8	When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison testing implemented. The results are recorded and this record is maintained.

K	2, 11		2.7.9	New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2		2.7.10	The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2		2.7.11	Membrane filters which are beyond their expiration date are not used.
O	11		2.7.12	Forceps tips are clean.
O	11		2.7.13	Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.
K	11		2.7.14	Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11		2.7.15	If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11		2.7.16	Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11		2.7.17	Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26		2.7.18	A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11		2.7.19	The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2		2.7.20	Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
2.8 Media Preparation and Storage – MF using mTEC Agar				
K	11		2.8.1	Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
C	11		2.8.2	The phosphate buffered saline is properly sterilized.
K	23		2.8.3	A sufficient amount of medium (4-5 mL) is used in each plate.
O	11		2.8.4	Refrigerated prepared plates are stored for no more than 2 weeks in sealed plastic bags or containers to minimize evaporation.
2.9 Sample Analyses - MF using mTEC Agar				
C	24		2.9.1	mTEC agar is used.
C	2		2.9.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
C	23, 35		2.9.3	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration.
C	23		2.9.4	The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25		2.9.5	Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23		2.9.6	Sample volumes are filtered under vacuum.
K	26		2.9.7	The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26		2.9.8	The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.
C	23		2.9.9	The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11		2.9.10	Blanks are run at the beginning of filtration, after every 10th aliquot and at the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).

C	2, 11	2.9.11	Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24	2.9.12	Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at 35 +0.5°C for 2 hours of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5°C for 24 ± 2 hours.
C	11, 23, 24	2.9.13	After 2 hours of resuscitation at 35°C, the watertight, tightly sealed containers are transferred to a circulating waterbath at 44.5 + 0.2°C, submerged completely and incubated for 22-24 hours.
			2.10 Computation of Results - MF using mTEC Agar
C	23	2.10.1	All yellow, yellow-green or yellow-brown colonies are counted.
C	23	2.10.2	Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as >80 x 100/the volume of sample filtered.
C	2, 11, 23	2.10.3	When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11	2.10.4	The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11	2.10.5	Results are reported as CFU/100 mL of sample.
			2.11 Bacteriological Analysis of UV Treated Process Water Samples by Membrane Filtration (MF) using mEndo Agar LES – Materials and Equipment
C	9, 11, 21	2.11.1	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches are used.
C	2	2.11.2	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
C	11, 19, 21	2.11.3	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm diameter with a pore size of 0.45µm and certified by the manufacturer for total coliform analysis.
C	2	2.11.4	Membrane filter lot numbers, dates of receipt and expiration dates are recorded and the records maintained.
C	2	2.11.5	If previous lots of agar or membrane filters are not available for comparability testing, an appropriate method for determining lot suitability has been developed and comparison testing implemented when the following has occurred: a. initiating monitoring by mEndo Agar LES; b. changing agar manufacturers; or c. changing brands of membrane filters used. The results are recorded and the records are maintained.
K	2, 11	2.11.6	Recovery of total coliforms from new lots of membrane filters and mEndo Agar LES is compared against the recovery from the previously acceptable lot.
C	2	2.11.7	The sterility of each lot and autoclave batch of membrane filters is verified before use.
K	2	2.11.8	Expired membrane filters are not used.
K	9, 11, 19, 21	2.11.9	Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.

K	11		2.11.10 If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
C	9, 11		2.11.11 Membrane filter assemblies are autoclave sterilized for 15 minutes at 121+/-2°C prior to the start of a filtration run. A new run occurs when there is a break of 30 minutes or more between the previous filtration run.
O	11, 19, 26, 36		2.11.12 A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11		2.11.13 The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and the records are maintained.
K	2		2.11.14 Maintenance of the UV sterilization unit is performed as needed. Maintenance is documented and the records maintained.
O	9, 11		2.11.15 Forceps tips are clean and smooth without pitting or corrugations.
			2.12 Media Preparation and Storage
C	9, 11, 19, 21, 36		2.12.1 mEndo Agar LES is used.
K	11, 21, 36		2.12.2 mEndo Agar LES is prepared aseptically using pre-sterilized glassware, sterile reagent water and pre-sterilized stir bar.
K	9, 11, 36		2.12.3 mEndo Agar LES is prepared using 95% alcohol that is not denatured.
C	9, 11, 36		2.12.4 mEndo Agar LES is brought to near boiling; removed from the heat and tempered at 44-50°C before dispensing.
C	9, 11, 36		2.12.5 mEndo Agar LES is never autoclaved.
K	9, 11, 36		2.12.6 A sufficient amount of medium (4-5 mL) is aseptically aliquoted to each culture plate.
O	9, 11, 36		2.12.7 Prepared plates of mEndo Agar LES are stored at 4°C in the dark for no more than two (2) weeks in sealed plastic bags or containers to minimize evaporation.
C	2		2.12.8 Appropriate, properly diluted positive and negative productivity controls for mEndo Agar LES medium are used. Results are recorded and the record maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11, 21, 36		2.12.9 Sterile phosphate buffered water or sterile phosphate buffered saline is used as a sample blank, filter funnel rinse and process and productivity control diluent for UV treated process water samples.
C	11		2.12.10 The phosphate buffered water/saline is properly sterilized and the sterility is tested before being placed in service. Results are recorded and records maintained
			2.13 Sample Analysis
C	9, 11, 36		2.13.1 The sample is shaken vigorously (25 times in a 12-inch arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration.
C	11, 21, 36		2.13.2 The membrane filter is placed grid side up within the sterile filter apparatus.
C	11, 26, 36		2.13.3 A 100 mL quantity of sample is filtered under vacuum.
K	26		2.13.4 The pressure of the vacuum pump does not exceed 15 psi.
C	9, 11, 26, 36		2.13.5 The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered water/saline as appropriate after filtration.
C	9, 11, 36		2.13.6 The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mEndo Agar LES so that no bubbles form between the filter and the agar.

K	9, 11, 36		2.13.7 Forceps are dipped in alcohol and flame sterilized between sample filters.
C	11, 36		2.13.8 Blanks are run at the beginning and at the end of the filtration run to check the sterility of the testing system (phosphate buffered water/saline, filter funnels, forceps, membrane filters, media and culture plates).
C	2, 36		2.13.9 An appropriate properly diluted positive process control culture accompanies the sample throughout incubation. Results are recorded and the records are maintained. Positive process control _____
C	9, 11, 36		2.13.10 Inoculated plates are incubated inverted at 35+/- 0.5°C for 22 to 24 hours.
K	2, 9, 11		2.13.11 An appropriate level of humidity is maintained in the incubator to prevent the plates from drying out.
2.14 Computation of Results			
K	9, 11		2.14.1 Colonies are counted with the aid of magnification.
C	9, 11, 19, 21, 23		2.14.2 All metallic sheen colonies are counted as total coliforms.
C	9, 11, 21, 36		2.14.3 Results are reported as total coliforms/100mL.
C	11, 20, 36		2.14.4 When no colonies are observed, results are reported as <1.0 coliform/100mL (nondetectable)
PART III - SHELLFISH SAMPLES			
3.1 Collection and Transportation of Samples			
C	9		3.1.1 A representative sample of shellstock is collected.
K	9		3.1.2 Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9		3.1.3 Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9		3.1.4 Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1		3.1.5 Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
3.2 Preparation of Shellfish for Examination			
K	2,11		3.2.1 Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2		3.2.2 Blades of shucking knives are not corroded.
O	9		3.2.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2		3.2.4 The faucet used for rinsing the shellstock does not contain an aerator.
K	9		3.2.5 Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9		3.2.6 Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	1, 9		3.2.7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9		3.2.8 Shellstock are not shucked directly through the hinge.
C	9		3.2.9 Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9		3.2.10 At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.

K	9		3.2.11 A representative sample of at least 12 shellfish is used for the analysis.
K	2		3.2.12 The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
O	9		3.2.13 Sterile phosphate buffered dilution water is used as the sample diluent.
C	9		3.2.14 Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9		3.2.15 APHA <i>Recommended Procedures for the Examination of Sea Water And Shellfish</i> , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA			
C	9		3.3.1 Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. (Circle the medium used.)
C	2		3.3.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9		3.3.3 Immediately (within 2 minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9		3.3.4 No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
C	9		3.3.5 Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6		3.3.6 In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2		3.3.7 Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9		3.3.8 Inoculated media are incubated at $35 \pm 0.5^\circ\text{C}$.
K	10		3.3.9 Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA			
C	9		3.4.1 EC medium is used as the confirmatory medium.
C	2		3.4.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control _____ Negative productivity control _____
K	9, 11		3.4.3 Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. (Circle the method of transfer.)
C	9		3.4.4 EC tubes are incubated in a circulating waterbath at $44.5 \pm 0.2^\circ\text{C}$
K	9		3.4.5 EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9		3.4.6 The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.
3.5 Computation of Results for MPN Analyses			
K	9		3.5.1 Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.

K	7		3.5.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9		3.5.3	Results are reported as MPN/100 grams of sample.
			3.6 Standard Plate Count Method	
O	20		3.6.1	A standard plate count (SPC) analysis maybe performed in conjunction with the analysis for fecal coliform organisms.
K	9		3.6.2	In the standard plate count procedure at least four plates are used, duplicates of two dilutions. One of the dilutions should produce colonies of 30 to 300 per plate.
K	2		3.6.3	Fifteen to 20 mL of tempered sterile plate count agar is used per plate.
C	9		3.6.4	Agar tempering bath maintains the agar at 44-46°C.
C	9		3.6.5	An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9		3.6.6	Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in 7 seconds) before plating.
C	9		3.6.7	Not more than 1 mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11		3.6.8	Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9,21		3.6.9	Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four high.
K	9		3.6.10	Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1		3.6.11	A hand tally or its equivalent is used for accuracy in counting.
			3.7 Computation of Results -SPC	
K	9		3.7.1	Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19		3.7.2	Colony counts are reported as CFU/g of sample.
			3.8 Bacteriological Analysis of Shellfish Using the ETCP	
C	2,3		3.8.1	Prepared modified MacConkey agar is used on the day that it is made.
K	3		3.8.2	Double strength modified MacConkey agar is used.
C	3		3.8.3	Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3		3.8.4	Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50°C until used.
K	2, 3		3.8.5	Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3		3.8.6	The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9		3.8.7	The sample homogenate is cultured within 2 minutes of blending.
C	2,3		3.8.8	Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.
K	3		3.8.9	Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2,3, 22		3.8.10	The container is gently swirled or slowly inverted once to mix the contents, which are subsequently distributed uniformly over six plates.
C	1		3.8.11	Media and diluent sterility are determined with each use. Results are recorded and the records maintained.

C	1		3.8.12 Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture _____ Negative control culture _____
C	3, 13		3.8.13 When solidified, the plates are placed inverted into an air incubator at 45.5 ± 0.5°C for 18 to 30 hours of incubation.
C	2		3.8.14 Plates are stacked no more than three high in the incubator.
C	2		3.8.15 Appropriately diluted pour plated process control cultures accompany each set of samples throughout incubation. The results are recorded and the records maintained. Positive process control _____ Negative process control _____
			3.9 Computation of Results - ETCP
K	11		3.9.1 Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1		3.9.2 A hand tally or its equivalent is used to aid in counting.
C	3, 6		3.9.3 All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3		3.9.4 Results are reported as CFU/100 grams of sample.
			Bacteriological Examination of Soft-shelled Clams and American Oysters for Male Specific Coliphage (MSC)
			3.10 MSC Equipment and Supplies
K	30		3.10.1 Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold 100 – 125 mL.
C	27, 28		3.10.2 The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	9		3.10.3 The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
C	27, 28		3.10.4 Sterile 0.22 µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1		3.10.5 The sterility of each lot of pre-sterilized syringes and syringe filters is determined. Results are recorded and records maintained.
K	1		3.10.6 The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28		3.10.7 The balance used provides a sensitivity of at least mg (0.01g.).
C	27, 28		3.10.8 The temperature of the incubator used is maintained at 36 ± 1°C.
C	28		3.10.9 Sterile disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
			3.11 MSC Media Preparation
K	28		3.11.1 Media preparation and sterilization is according to the validated method.
K	27, 28		3.11.2 Bottom agar, double strength soft agar and growth broth are prepared from their individual components.
K	27, 28		3.11.3 Soft agar is prepared double strength in volumes of 2.5 mL.
C	27, 28		3.11.4 The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
O	27, 28		3.11.5 Storage of the bottom agar under refrigeration does not exceed 1 month.
K	27, 28		3.11.6 Unsterilized soft agar is stored at -20 °C -15C for up to 3 months.
K	27, 28		3.11.7 The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.
K	27, 28		3.11.8 Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	27, 28		3.11.9 Bottom agar plates are allowed to reach room temperature before use.

3.12 Preparation of the Soft-Shelled Clams and American Oysters for MSC Analysis		
K	2,11	3.12.1 Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2	3.12.2 The blades of shucking knives are not corroded.
O	9	3.12.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2	3.12.4 The faucet used for rinsing the shellfish does not contain an aerator.
K	9	3.12.5 The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9	3.12.6 The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9	3.12.7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9	3.12.8 Shellfish are not shucked through the hinge.
C	9	3.12.9 The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9	3.12.10 A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19	3.12.11 The sample is weighed to the nearest 0.1 gram.
3.13 MSC Sample Analysis		
C	28	3.13.1 <i>E. coli</i> Famp ATCC 700891 is the bacterial host strain used in this procedure.
K	27, 28	3.13.2 Host cell growth broth is tempered at $36 \pm 1^\circ\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	27, 28	3.13.3 Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^\circ\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	27, 28	3.13.4 After inoculation, the host cell growth broth culture is not shaken.
C	28	3.13.5 A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.
C	28	3.13.6 The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28	3.13.7 The elution mixture is homogenized at high speed for 180 seconds.
C	28	3.13.8 Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
C	28	3.13.9 The homogenized elution mixture is centrifuged for 15 minutes at $9000 \times g$ at 4°C.
C	27, 28	3.13.10 The supernatant is pipetted off, weighed and the weight recorded.
C	27, 28	3.13.11 The supernatant is allowed to warm to room temperature about 20 to 30 minutes.
K	27, 28	3.13.12 The autoclaved soft agar is tempered and held at $51 \pm 1^\circ\text{C}$ throughout the period of sample analysis.
K	27, 28	3.13.13 Two hundred microliters (0.2 mL) of log phase host strain <i>E. coli</i> is added to the tempering soft agar immediately prior to adding the sample supernatant.
K	27, 28	3.13.14 The sample supernatant is shaken or vortexed before being added to the tempering soft agar.
C	27, 28	3.13.15 2.5 mL of sample supernatant is added to each tube of tempering soft agar.
C	27, 28	3.13.16 The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28	3.13.17 The soft agar/sample supernatant/host cell mixture is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.
C	28	3.13.18 Ten (10) plates are used, 2.5 mL per plate for a total of 25 mL of supernatant analyzed per sample.

K	27, 28		3.13.19 Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are recorded and records maintained. Positive control _____
K	27, 28		3.13.20 Growth broth is used as the negative control or blank.
K	27, 28		3.13.21 Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2		3.13.22 A negative control plate is plated at the beginning and end of each set of samples analyzed.
K	27, 28		3.13.23 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28		3.13.24 All plates are incubated at 36 ± 1°C for 18 ± 2 hours.
3.14 Computation of Results - MSC			
C	27		3.14.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 32		3.14.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten plates, the count is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/ 100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as > 20,000 PFU/100 grams.
K	28		3.14.3 The formula used for determining the density of MSC in PFU/100 grams is: (0.364) (N) (Ws), where N = total number of plaques counted on all 10 plates and Ws = weight of the supernatant used.
O	9		3.14.4 The MSC count is rounded off conventionally to give a whole number.

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LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
MICROBIOLOGICAL COMPONENT: (Part I-III)	
A. Results	
Total # of Critical (C) Nonconformities in Parts I-III	_____
Total # of Key (K) Nonconformities in Parts I-III	_____
Total # of Critical, Key and Other (O) Nonconformities in Parts I-III	_____
B. Criteria for Determining Laboratory Status of the Microbiological Component:	
<p>1. Does Not Conform Status: The Microbiological component of this laboratory is not in conformity with NSSP requirements if:</p> <ul style="list-style-type: none"> a. The total # of Critical nonconformities is ≥ 4 or b. The total # of Key nonconformities is ≥ 13 or c. The total # of Critical, Key and Other is ≥ 18 <p>2. Provisionally Conforms Status: The microbiological component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but ≤ 3.</p>	
C. Laboratory Status (circle appropriate)	
Does Not Conform	Provisionally Conforms
Conforms	
<p>Acknowledgment by Laboratory Director/Supervisor:</p> <p>All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____.</p> <p>Laboratory Signature: _____ Date: _____</p> <p>LEO Signature: _____ Date: _____</p>	

PART 1 - QUALITY ASSURANCE		
CODE	REF.	ITEM
K	8, 11	1.1 Quality Assurance (QA) Plan
		1.1.1 Written Plan (Check those items which apply.)
		a. Organization of the laboratory.
		b. Staff training requirements.
		c. Standard operating procedures.
		d. Internal quality control measures for equipment, their calibration, maintenance, repair, performance, and rejection criteria established.
		e. Laboratory safety.
		f. Internal performance assessment.
		g. External performance assessment.
C	8	1.1.2 QA Plan Implemented.
K	11	1.1.3 The Laboratory participates in a proficiency testing program annually. Specify Program(s)_____
		1.2 Educational/Experience Requirements
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, or equivalent discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial laboratories, the analyst(s) must have at least a high school diploma and shall have at least three months of experience in laboratory science
		1.3 Work Area
O	8,11	1.3.1 Adequate for workload and storage.
K	11	1.3.2 Clean, well-lighted.
K	11	1.3.3 Adequate temperature control.
O	11	1.3.4 All work surfaces are nonporous, easily cleaned and disinfected.
K	11	1.3.5 Microbiological quality of the air is fewer than 15 colonies for a 15 minute exposure and determined monthly. The results are recorded and records maintained.
		1.4 Laboratory Equipment
O	9	1.4.1 To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.
O	14	1.4.2 pH electrodes consisting of pH half-cell and reference half-cell or equivalent combination electrode free from (Ag/AgCl) or contains an ion exchange barrier preventing passage of Ag ions into the medium which may affect the accuracy of the pH reading.
K	11	1.4.3 The effect of temperature on the pH is compensated for by an ATC probe or by manual adjustment.
K	8	1.4.4 pH meter is calibrated daily or with each use Results are recorded and records maintained.
K	11	1.4.5 A minimum of two standard buffer solutions is used to calibrate the pH meter. The first must be near the electrode isopotential point (pH 7). The second near the expected sample pH (i.e., pH 4 or pH 10). Standard buffer solutions are used once and discarded.
O	8,15	1.4.6 Electrode acceptability is determined daily or with each use by the millivolt procedure or through determination of the slope. (Circle the method used.)

K	9		1.4.7 Balance provides a sensitivity of at least 0.1 g at weights of use.
K	11,13		1.4.8 Balance calibrations are checked monthly according to manufacturer's specifications using NIST Class S or ASTM Class 1 or 2 weights or equivalent. The accuracy of the balance is verified at the weight range of use. Results are recorded and records maintained.
K	11		1.4.9 Refrigerator temperature(s) are monitored at least once daily on workdays Results are recorded and records maintained.
K	1		1.4.10 Refrigerator temperature is maintained at 0 to 4°C.
C	9		1.4.11 The temperature of the incubator is maintained at 35 ± 0.5°C.
C	11		1.4.12 Thermometers used in the air incubator(s) are graduated in at least 0.1°C increments.
K	9		1.4.13 Working thermometers are located on top and bottom shelves or appropriately placed based on the results of spatial temperature checks.
C	11		1.4.14 Temperature of the waterbath is maintained at 44.5 ± 0.2°C under all loading conditions.
C	9		1.4.15 The thermometers used in the waterbath are graduated in at least 0.1°C increments.
C	13		1.4.16 The waterbath has adequate capacity for workload.
K	9		1.4.17 The level of water in the waterbath covers the level of liquid in the incubating tubes.
K	8, 11		1.4.18 Air incubator/waterbath temperatures are taken twice daily on workdays. The results are recorded and records maintained.
C	4		1.4.19 All working thermometers are appropriately immersed.
C	29		1.4.20 Working thermometers are either: calibrated mercury-in-glass thermometers, calibrated non-mercury-in-glass thermometers, or appropriately calibrated electronic devices, including Resistance Temperature Devices (RTDs) and Platinum Resistance Devices (PTDs).
C	11		1.4.21 A standards thermometer has been calibrated by NIST or a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at the points 0, 35 and 44.5°C (45.5°C for ETCP). These calibration records are maintained.
K	9		1.4.22 Standards thermometers are checked annually for accuracy by ice point determination. Results recorded and maintained. Date of most recent determination_____.
C	29		1.4.23 Either mercury-in-glass thermometers, non-mercury-in-glass thermometers having the accuracy (uncertainty), tolerance and response time of mercury or low drift electronic resistance thermometers with an accuracy of ≤ ±0.05°C are used as the laboratory standards thermometer. (Circle the thermometer type used.)
K	13		1.4.24 The accuracy of working thermometers is checked annually against the standards thermometer either at the temperatures at which they are used or by ice point determination. Results are recorded and records maintained.
O	11		1.4.25 Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
1.5 Labware and Glassware Washing			
O	9		1.5.1 Utensils and containers are clean borosilicate glass, stainless steel or other noncorroding materials.
K	9		1.5.2 Culture tubes are of a suitable size to accommodate the volume for nutritive ingredients and samples.
K	9		1.5.3 Sample containers are made of glass or some other inert material.
O	9		1.5.4 Dilution bottles and tubes are made of borosilicate glass or plastic and closed with rubber stoppers, caps or screw caps with nontoxic liners.

K	9		1.5.5	Graduations are indelibly marked on dilution bottles and tubes or an acceptable alternative method is used to ensure appropriate volumes.
C	9		1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken tips and are appropriately graduated. Pipettes larger than 10 mL are not used to deliver 1mL aliquots; nor, are pipets larger than 1.1mL used to deliver 0.1 mL aliquots.
K	9		1.5.7	Reusable sample containers are capable of being properly washed and sterilized.
K	9		1.5.8	In washing reusable pipettes, a succession of at least three fresh water rinses plus a final rinse of distilled/deionized water is used to thoroughly rinse off all the detergent.
C	2		1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
C	11		1.5.10	With each load of labware/glassware washed the contact surface of several dry pieces from each load are tested for residual detergent (acid or alkali) with aqueous 0.04% bromothymol blue. Results are recorded and records maintained.
1.6 Sterilization and Decontamination				
K	9		1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
O	8		1.6.2	Routine autoclave maintenance is performed and the records are maintained.
C	30, 33, 34		1.6.3	The autoclave provides sterilization conditions suitable to the load contents. Sterilization temperature range may be 119°C - 124°C as determined by the lab's equipment Quality Assurance Verification Testing and recommended practices from the media manufacturer. Sterilization is determined for each load using a working maximum registering thermometer or an appropriate working temperature monitoring device.
K	11		1.6.4	An autoclave standards thermometer has been calibrated by a qualified calibration laboratory using a primary standard traceable to NIST or an equivalent authority at 121°C. Calibration at 100°C, the steam point, is also recommended but not required.
K	16		1.6.5	The autoclave standards thermometer is checked every five (5) years for accuracy at 121°C by a qualified calibration laboratory; or, is checked in-house at the steam point (100°C) if it has been previously calibrated at both 100°C and 121°C. Any change in temperature at the steam point changes the calibrated temperature at 121°C by the same magnitude. Date of most recent determination _____
K	1		1.6.6	Working autoclave thermometers are checked against the autoclave standards thermometer at 121°C yearly. Date of last check _____ Method _____
K	11		1.6.7	Spore strips/suspensions appropriate for use in an autoclave media cycle are used monthly according to manufacturer's instructions to evaluate the effectiveness of the sterilization process. Results are recorded and the records maintained.
O	11		1.6.8	Heat sensitive tape is used with each autoclave batch.
K	11, 13		1.6.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained. Type of record: Autoclave log, computer printout or chart recorder tracings. (Circle appropriate type or types.)
K	11		1.6.10	For dry heat sterilized material, the hot-air sterilizing oven provides heating and sterilizing temperatures in the range of 160 to 180°C.
K	9		1.6.11	A thermometer capable of determining temperatures accurately in the range of 160 to 180°C is used to monitor the operation of the hot-air sterilizing oven
K	13		1.6.12	Records of temperatures and exposure times are maintained for the operation of the hot-air sterilizing oven during use.

K	11		1.6.13 Spore strips/suspensions are used quarterly to evaluate the effectiveness of the sterilization process in the hot-air oven. Records are maintained.
K	11		1.6.14 Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air oven or autoclaved for 15 minutes at 121°C.
C	1		1.6.15 The sterility of reusable sample containers is determined for each load sterilized. The results are recorded and the records maintained.
C	1		1.6.16 The sterility of pre-sterilized disposable sample containers is determined for each lot received. Results are recorded and the records maintained.
K	9		1.6.17 Reusable pipettes are stored and sterilized in aluminum or stainless steel canisters.
K	9		1.6.18 Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2 hours.
C	2		1.6.19 The sterility of reusable pipettes is determined with each load sterilized. Results are recorded and records maintained.
C	2		1.6.20 The sterility of pre-sterilized disposable pipettes is determined with each lot received. Results are recorded and the records maintained.
K	18		1.6.21 Hardwood applicator transfer sticks are properly sterilized. Method of sterilization _____
C	2		1.6.22 The sterility of the hardwood applicator transfer sticks is checked routinely. Results are recorded and the records maintained.
O	13		1.6.23 Spent broth cultures and agar plates are decontaminated by autoclaving for at least 30 minutes before conventional disposal.
1.7 Media Preparation			
K	3, 5		1.7.1 Media is commercially dehydrated except in the case of medium A-1 which must be prepared from the individual components and modified MacConkey agar which may be prepared from its components.
K	11		1.7.2 Media is prepared according to manufacturer's instructions.
O	11		1.7.3 Dehydrated media and media components are properly stored in a cool, clean, dry place.
O	11		1.7.4 Dehydrated media are labeled with date of receipt and date opened.
C	12		1.7.5 Caked or expired media or media components are discarded.
C	11		1.7.6 Reagent water is distilled or deionized (circle appropriate choice), tested monthly and exceeds 0.5 megohm-cm resistance (2 megohms-cm in-line) or is less than 2.0 µSiemens/cm conductivity at 25°C. (Circle the appropriate water quality descriptor determined.) Results are recorded and the records maintained.
C	11		1.7.7 Reagent water is analyzed for residual chlorine monthly and is at a non-detectable level (<0.1 mg/L). Results are recorded and the records maintained. Specify method of determination_____.
K	11		1.7.8 Reagent water contains <100 CFU/mL as determined monthly using the heterotrophic plate count method. Results are recorded and the records maintained.
K	11		1.7.9 Media prepared from commercial dehydrated components are sterilized according to the manufacturer's instructions.
K	9		1.7.10 The volume and concentration of media in the tube are suitable for the amount of sample inoculated.
C	11		1.7.11 Total time of exposure of sugar broths to autoclave temperatures does not exceed 45 minutes.
C	1		1.7.12 Media sterility is determined for each load sterilized. Results are recorded and the records maintained.

C	1	1.7.13 Media productivity is determined using media-appropriate, properly diluted positive and negative control cultures for each lot of dehydrated media received or with each batch of media prepared when the medium is made from its individual components.
		Media: Positive control: Negative control:
		Media: Positive control: Negative control:
		Media: Positive control: Negative control:
		Media: Positive control: Negative control:
		Media: Positive control: Negative control:
		Media: Positive control: Negative control:
O	9	1.7.14 Sterile phosphate buffered dilution water is used as the sample diluent.
K	11	1.7.15 The pH of the prepared media is determined after sterilization to ensure that it is consistent with manufacturer's requirements. Results are recorded and records are maintained.
1.8 Storage of Prepared Culture Media		
K	9	1.8.1 Prepared culture media are stored in a cool, clean, dry space where excessive evaporation and the danger of contamination are minimized.
K	5,11	1.8.2 Brilliant green bile 2% broth and A-1 media are stored in the dark.
K	13	1.8.3 Stored media are labeled with the storage expiration date or the sterilization date.
K	9	1.8.4 Storage of prepared culture media at room temperature does not exceed 7 days.
K	2	1.8.5 Storage under refrigeration of prepared culture media with loose fitting closures shall not exceed 1 month.
K	11	1.8.6 Storage under refrigeration of prepared culture media with screw-cap closures does not exceed 3 months.
K	17	1.8.7 All prepared MPN broth media stored under refrigeration must reach room temperature prior to use. Culture tubes containing any type of precipitate or Durham tubes containing air bubbles are discarded.
PART II - SEAWATER SAMPLES		
2.1 Collection and Transportation of Samples		
C	11	2.1.1 Sample containers are of a suitable size to contain at least 110 mL of sample and to allow adequate headspace for proper shaking. Seawater samples are collected in clean, sterile, watertight, properly labeled sample containers.
K	1	2.1.2 Samples are identified with collector's name, harvest area, sampling station, time and date of collection.
C	9	2.1.3 Immediately after collection, seawater samples are placed in dry storage (ice chest or equivalent) capable of maintaining a temperature of 0 to 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed in the refrigerator unless processed immediately.
O	1	2.1.4 A temperature blank is used to represent the temperature of samples upon receipt at the laboratory. Temperature should be equivalent or less than that of the growing area waters. Results are recorded and maintained.
C	9	2.1.5 Analysis of the sample is initiated as soon as possible after collection. Seawater samples are not tested if they have been held for more than 30 hours from the time of collection.

			2.2 Bacteriological Examination of Seawater by the APHA MPN
C	9		2.2.1 Lactose broth or lauryl tryptose broth is used as the presumptive medium. <i>(Circle appropriate one.)</i>
€	2		2.2.2 The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control Negative productivity control
C	9, 35		2.2.3 2 Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9		2.2.4 3 In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6		2.2.5 4 In a single dilution series not less than 12 tubes are used (for depuration at least 5 tubes are used).

C	6	2.2.65	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
K	9	2.2.76	Inoculated tubes are incubated in air at 35 ± 0.5°C.
C	2	2.2.87	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
K	9	2.2.98	Inoculated tubes are read after 24 ± 2 hours and 48 ± 3 hours of incubation and transferred at both time interval if positive for growth (the presence of turbidity) and gas or effervescence in the culture tube. These tubes are considered presumptive positive requiring further confirmatory testing.
2.3 Confirmed Test for Seawater by APHA MPN			
C	9	2.3.1	Brilliant green bile 2% broth (BGB) is used as the confirmatory medium for total coliforms.
C	9	2.3.2	EC medium is used as the confirmatory medium for fecal coliforms.
€	2	2.3.3	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control Negative productivity control
K	9, 11	2.3.43	Transfers are made to BGB/EC by either sterile loop or sterile hardwood transfer stick from positive presumptive tubes incubated for 24 and 48 hours as appropriate. (Circle the method of transfer.)
C	9	2.3.54	BGB tubes are incubated at 35 ± 0.5°C.
K	9	2.3.65	BGB tubes are read after 48 ± 3 hours of incubation.
C	9	2.3.76	EC tubes are incubated in a circulating waterbath maintained at 44.5 ± 0.2°C.
C	9	2.3.87	EC tubes are read after 24 ± 2 hours of incubation.
C	9	2.3.98	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.4 Computation of Results – APHA MPN			
K	9	2.4.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
K	7	2.4.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.4.3	Results are reported as MPN/100 mL of sample.
2.5 Bacteriological Examination of Seawater by the MA-1 Method			
C	5	2.5.1	A-1 medium complete is used in the analysis.
C	2, 31	2.5.2	A-1 medium without salicin is used in the analysis. Comparability testing supports use of A-1 medium without salicin. Study records are available.
C	5	2.5.3	A-1 medium sterilized for 10 minutes at 121°C.
€	2	2.5.4	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control Negative productivity control

C	9, 35	2.5.54	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before inoculation.
C	9	2.5.65	In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes are recommended).
C	6	2.5.76	In a single dilution series at least 12 tubes are used.
C	6	2.5.87	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	2.5.98	Appropriately diluted process control cultures accompany the samples <i>throughout both resuscitation and waterbath incubation</i> Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	2,5	2.5.109	Inoculated tubes are placed in an air incubator at $35 \pm 0.5^\circ\text{C}$ for 3 ± 0.5 hours of resuscitation.
C	5	2.5.110	After 3 ± 0.5 hours resuscitation at 35°C , inoculated tubes are incubated at $44.5 \pm 0.2^\circ\text{C}$ in a circulating waterbath for the remainder of the 24 ± 2 hours.
C	5	2.5.121	The presence of turbidity and any amount of gas or effervescence in the culture tube constitutes a positive test.
2.6 Computation of Results – APHA MPN			
K	9	2.6.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition.
K	7	2.6.2	Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	7, 9	2.6.3	Results are reported as MPN/100 mL of sample.
2.7 Bacteriological Analysis of Seawater by Membrane Filtration (MF) using mTEC Agar - Materials and Equipment			
C	23, 24	2.7.1	When used for elevated temperature incubation in conjunction with ethafoam resuscitation, the temperature of the hot air incubator is maintained at $44.5 \pm 0.5^\circ\text{C}$ under any loading capacity.
C	23	2.7.2	When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
C	23	2.7.3	Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches with tight fitting lids are used.
C	2	2.7.4	The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
K	11	2.7.5	Colonies are counted with the aid of magnification.
C	11, 23	2.7.6	Membrane filters are made from cellulose ester material, white, grid marked, 47 mm in diameter with a pore size of $0.45 \mu\text{m}$ and certified by the manufacturer for fecal coliform analyses.
C	2	2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
C	2	2.7.8	When initiating monitoring by mTEC or switching brands or types of membrane filters used and no previous lots of filters are available for comparing acceptable performance, an appropriate method for determining the suitability of the lot is developed and the comparison testing implemented. The results are recorded and this record is maintained.

K	2, 11	2.7.9	New lots of membrane filters are checked by comparing recovery of fecal coliform organisms against membrane filters from previously acceptable lots.
C	2	2.7.10	The sterility of each lot or autoclave batch of membrane filters are checked before use.
K	2	2.7.11	Membrane filters which are beyond their expiration date are not used.
O	11	2.7.12	Forceps tips are clean.
O	11	2.7.13	Forceps tips are smooth without pitting or corrugations to damage the filters being manipulated.
K	11	2.7.14	Forceps are dipped in alcohol and flame sterilized between sample filters.
K	11	2.7.15	If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically or with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
K	11	2.7.16	Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.
C	11	2.7.17	Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
O	11, 23, 26	2.7.18	A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11	2.7.19	The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and records maintained.
K	2	2.7.20	Maintenance of the UV sterilization unit is performed as needed. This maintenance is documented and the records maintained.
2.8 Media Preparation and Storage – MF using mTEC Agar			
K	11	2.8.1	Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
C	11	2.8.2	The phosphate buffered saline is properly sterilized.
K	23	2.8.3	A sufficient amount of medium (4-5 mL) is used in each plate.
O	11	2.8.4	Refrigerated prepared plates are stored for no more than 2 weeks in sealed plastic bags or containers to minimize evaporation.
2.9 Sample Analyses - MF using mTEC Agar			
C	24	2.9.1	mTEC agar is used.
e	2	2.9.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained.
C	23, 35	2.9.3	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration.
C	23	2.9.4	The membrane is placed grid side up within the sterile filter apparatus.
C	23, 25	2.9.5	Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling).
C	23	2.9.6	Sample volumes are filtered under vacuum.
K	26	2.9.7	The pressure of the vacuum pump does not exceed 15 psi.
C	23, 26	2.9.8	The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration.
C	23	2.9.9	The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mTEC agar so that no bubbles form between the filter and the agar.
C	11	2.9.10	Blanks are run at the beginning of filtration, after every 10th aliquot and at the end of the filtration run to check the sterility of the testing system (phosphate buffered saline, filter funnel, forceps, membrane filter, media and culture plate).

C	2, 11		2.9.110 Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are recorded and the records maintained. Positive process control _____ Negative process control _____
C	11, 23, 24		2.9.121 Inoculated plates are placed inverted into a watertight, tightly sealed container prior to being placed in the air incubator and incubated at 35 + 0.5°C for 2 hours of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5°C for 24 ± 2 hours.
C	11, 23, 24		2.9.132 After 2 hours of resuscitation at 35°C, the watertight, tightly sealed containers are transferred to a circulating waterbath at 44.5 + 0.2°C, submerged completely and incubated for 22-24 hours.
		2.10 Computation of Results - MF using mTEC Agar	
C	23		2.10.1 All yellow, yellow-green or yellow-brown colonies are counted.
C	23		2.10.2 Only plates having 80 or fewer colonies are counted. If it is unavoidable to use plates having more than 80 colonies, counts are given as >80 x 100/the volume of sample filtered.
C	2, 11, 23		2.10.3 When multiple dilutions are filtered, the laboratory has developed a procedure for assessing the contribution of all positive dilutions to the final count.
C	23, 11		2.10.4 The number of fecal coliforms is calculated by the following equation: Number of fecal coliforms per 100 mL = [number of colonies counted per plate used in the count / volume (s) of sample filtered in ml] x 100.
C	23, 11		2.10.5 Results are reported as CFU/100 mL of sample.
		2.11 Bacteriological Analysis of UV Treated Process Water Samples by Membrane Filtration (MF) using mEndo Agar LES – Materials and Equipment	
C	9, 11, 21		2.11.1 Pre-sterilized plastic or sterile glass culture plates that are clear, flat bottomed, free of bubbles and scratches are used.
C	2		2.11.2 The sterility of pre-sterilized culture plates is determined for each lot received. Results are recorded and the records maintained.
C	11, 19, 21		2.11.3 Membrane filters are made from cellulose ester material, white, grid marked, 47 mm diameter with a pore size of 0.45µm and certified by the manufacturer for total coliform analysis.
C	2		2.11.4 Membrane filter lot numbers, dates of receipt and expiration dates are recorded and the records maintained.
C	2		2.11.5 If previous lots of agar or membrane filters are not available for comparability testing, an appropriate method for determining lot suitability has been developed and comparison testing implemented when the following has occurred: a. initiating monitoring by mEndo Agar LES; b. changing agar manufacturers; or c. changing brands of membrane filters used. The results are recorded and the records are maintained.
K	2, 11		2.11.6 Recovery of total coliforms from new lots of membrane filters and mEndo Agar LES is compared against the recovery from the previously acceptable lot.
C	2		2.11.7 The sterility of each lot and autoclave batch of membrane filters is verified before use.
K	2		2.11.8 Expired membrane filters are not used.
K	9, 11, 19, 21		2.11.9 Membrane filtration units are made of stainless steel, glass or autoclavable plastic free of scratches, corrosion and leaks.

K	11		2.11.10 If indelible graduation marks are used on clear glass or plastic funnels to measure sample volumes, their accuracy is checked gravimetrically with a Class A graduated cylinder before use and periodically rechecked. Funnels having a tolerance greater than 2.5% are not used. Checks are recorded and records maintained.
C	9, 11		2.11.11 Membrane filter assemblies are autoclave sterilized for 15 minutes at 121+/- 2°C prior to the start of a filtration run. A new run occurs when there is a break of 30 minutes or more between the previous filtration run.
O	11, 19, 26, 36		2.11.12 A UV sterilization unit is used to disinfect filter assemblies between sample and filtration runs.
K	11		2.11.13 The effectiveness of the UV sterilization unit is determined by biological testing monthly. Results are recorded and the records are maintained.
K	2		2.11.14 Maintenance of the UV sterilization unit is performed as needed. Maintenance is documented and the records maintained.
O	9, 11		2.11.15 Forceps tips are clean and smooth without pitting or corrugations.
2.12 Media Preparation and Storage			
C	9, 11, 19, 21, 36		2.12.1 mEndo Agar LES is used.
K	11, 21, 36		2.12.2 mEndo Agar LES is prepared aseptically using pre-sterilized glassware, sterile reagent water and pre-sterilized stir bar.
K	9, 11, 36		2.12.3 mEndo Agar LES is prepared using 95% alcohol that is not denatured.
C	9, 11, 36		2.12.4 mEndo Agar LES is brought to near boiling; removed from the heat and tempered at 44-50°C before dispensing.
C	9, 11, 36		2.12.5 mEndo Agar LES is never autoclaved.
K	9, 11, 36		2.12.6 A sufficient amount of medium (4-5 mL) is aseptically aliquoted to each culture plate.
O	9, 11, 36		2.12.7 Prepared plates of mEndo Agar LES are stored at 4°C in the dark for no more than two (2) weeks in sealed plastic bags or containers to minimize evaporation.
€	2		2.12.8 Appropriate, properly diluted positive and negative productivity controls for mEndo Agar LES medium are used. Results are recorded and the record maintained. Positive productivity control Negative productivity control
K	9, 11, 21, 36		2.12.9 Sterile phosphate buffered water or sterile phosphate buffered saline is used as a sample blank, filter funnel rinse and process and productivity control diluent for UV treated process water samples.
C	11		2.12.10 The phosphate buffered water/saline is properly sterilized and the sterility is tested before being placed in service. Results are recorded and records maintained
2.13 Sample Analysis			
C	9, 11, 36		2.13.1 The sample is shaken vigorously (25 times in a 12-inch arc in 7 seconds by hand or for at least 15 seconds when using a mechanical shaker) before filtration.
C	11, 21, 36		2.13.2 The membrane filter is placed grid side up within the sterile filter apparatus.
C	11, 26, 36		2.13.3 A 100 mL quantity of sample is filtered under vacuum.
K	26		2.13.4 The pressure of the vacuum pump does not exceed 15 psi.
C	9, 11, 26, 36		2.13.5 The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered water/saline as appropriate after filtration.

C	9, 11, 36		2.13.6 The membrane filter is removed from the filtering apparatus with sterile forceps and rolled onto mEndo Agar LES so that no bubbles form between the filter and the agar.
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K	9, 11, 36		2.13.7 Forceps are dipped in alcohol and flame sterilized between sample filters.
C	11, 36		2.13.8 Blanks are run at the beginning and at the end of the filtration run to check the sterility of the testing system (phosphate buffered water/saline, filter funnels, forceps, membrane filters, media and culture plates).
C	2, 36		2.13.9 An appropriate properly diluted positive process control culture accompanies the sample throughout incubation. Results are recorded and the records are maintained. Positive process control _____
C	9, 11, 36		2.13.10 Inoculated plates are incubated inverted at 35+/- 0.5°C for 22 to 24 hours.
K	2, 9, 11		2.13.11 An appropriate level of humidity is maintained in the incubator to prevent the plates from drying out.
2.14 Computation of Results			
K	9, 11		2.14.1 Colonies are counted with the aid of magnification.
C	9, 11, 19, 21, 23		2.14.2 All metallic sheen colonies are counted as total coliforms.
C	9, 11, 21, 36		2.14.3 Results are reported as total coliforms/100mL.
C	11, 20, 36		2.14.4 When no colonies are observed, results are reported as <1.0 coliform/100mL (nondetectable)
PART III - SHELLFISH SAMPLES			
3.1 Collection and Transportation of Samples			
C	9		3.1.1 A representative sample of shellstock is collected.
K	9		3.1.2 Shellstock samples are collected in clean, waterproof, puncture resistant containers loosely sealed.
K	9		3.1.3 Shellstock samples are labeled with collector's name, type of shellstock, the source or harvest area, sampling station, time, date and place (if applicable) of collection.
C	9		3.1.4 Immediately after collection, shellfish samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10°C with ice or cold packs for transport to the laboratory. Once received, the samples are placed under refrigeration unless processed immediately.
C	1		3.1.5 Analysis of the samples is initiated as soon as possible after collection. Shellfish samples are not tested if the time interval between collection and analysis exceeds 24 hours.
3.2 Preparation of Shellfish for Examination			
K	2,11		3.2.1 Shucking knives, scrub brushes and blender jars are (autoclave) sterilized for 15 minutes prior to use.
O	2		3.2.2 Blades of shucking knives are not corroded.
O	9		3.2.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2		3.2.4 The faucet used for rinsing the shellstock does not contain an aerator.
K	9		3.2.5 Shellstock are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9		3.2.6 Shellstock are allowed to drain in a clean container or on clean towels prior to opening.
K	1, 9		3.2.7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9		3.2.8 Shellstock are not shucked directly through the hinge.
C	9		3.2.9 Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9		3.2.10 At least 200 grams of shellfish meat or a quantity of meat sufficient to cover the blender blades is used for the analysis.

K	9	3.2.11	A representative sample of at least 12 shellfish is used for the analysis.
K	2	3.2.12	The sample is weighed to the nearest 0.1 gram and an equal amount by weight of diluent is added.
O	9	3.2.13	Sterile phosphate buffered dilution water is used as the sample diluent.
C	9	3.2.14	Samples are blended at high speed for 60 to 120 seconds until homogenous.
K	9	3.2.15	APHA <i>Recommended Procedures for the Examination of Sea Water And Shellfish</i> , Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
3.3 MPN Analysis for Fecal Coliform Organisms, Presumptive Test, APHA			
C	9	3.3.1	Appropriate strength lactose or lauryl tryptose broth is used as presumptive media in the analysis. <i>(Circle the medium used.)</i>
€	2	3.3.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control Negative productivity control
K	9	3.3.32	Immediately (within 2 minutes) after blending, the ground sample is diluted and inoculated into tubes of presumptive media.
C	9	3.3.43	No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
C	9	3.3.54	Allowing for the initial 1:1 dilution of the sample, appropriate portions are inoculated (i.e., 2 ml of original 1:1 dilution for the 1 g portion) and diluted for subsequent inoculation (i.e., 20 ml of 1:1 diluted sample to 80 ml of diluent or the equivalent for 0.1 g portion). All successive dilutions are prepared conventionally.
K	6	3.3.65	In a single dilution series, the volumes examined are adequate to meet the needs of routine monitoring. Sample volume inoculated _____ Range of MPN _____ Strength of media used _____
C	2	3.3.76	Appropriately diluted process control cultures accompany the samples throughout both the presumptive and confirmed phases of incubation. Results are recorded and the records maintained. Positive Process control _____ Negative Process control _____
K	9	3.3.87	Inoculated media are incubated at $35 \pm 0.5^\circ\text{C}$.
K	10	3.3.98	Tubes are read after 24 ± 2 hours of incubation and transferred if positive for growth (the presence of turbidity and gas or effervescence in the culture tube). These tubes are considered presumptive requiring further confirmatory testing.
3.4 Confirmed Test for Fecal Coliforms - APHA			
C	9	3.4.1	EC medium is used as the confirmatory medium.
€	2	3.4.2	The appropriate positive and negative productivity controls for the presumptive media are used. The results are recorded and the records maintained. Positive productivity control Negative productivity control
K	9, 11	3.4.32	Transfers are made to EC medium by either sterile loop or hardwood sterile transfer sticks from positive presumptives. <i>(Circle the method of transfer.)</i>
C	9	3.4.43	EC tubes are incubated in a circulating waterbath at $44.5 \pm 0.2^\circ\text{C}$
K	9	3.4.54	EC tubes are read for gas production after 24 ± 2 hours of incubation.
C	9	3.4.65	The presence of turbidity and any amount of gas and/or effervescence in the Durham tube constitutes a positive test.
3.5 Computation of Results for MPN Analyses			
K	9	3.5.1	Results of multiple dilution tests are read from tables in <i>Recommended Procedure for the Examination of Sea Water and Shellfish</i> , 4th Edition and multiplied by the appropriate dilution factor.

K	7		3.5.2 Results from single dilution series are calculated from Hoskins' equation or interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube Method".
C	9		3.5.3 Results are reported as MPN/100 grams of sample.
			3.6 Standard Plate Count Method
O	20		3.6.1 A standard plate count (SPC) analysis maybe performed in conjunction with the analysis for fecal coliform organisms.
K	9		3.6.2 In the standard plate count procedure at least four plates are used, duplicates of two dilutions. One of the dilutions should produce colonies of 30 to 300 per plate.
K	2		3.6.3 Fifteen to 20 mL of tempered sterile plate count agar is used per plate.
C	9		3.6.4 Agar tempering bath maintains the agar at 44-46°C.
C	9		3.6.5 An agar based temperature control having a similar volume and shape as the tempering plate count agar is used in the tempering bath.
K	9		3.6.6 Samples or sample dilutions to be plated are shaken vigorously (25 times in a 12" arc in 7 seconds) before plating.
C	9		3.6.7 Not more than 1 mL nor less than 0.1 mL of sample or sample dilution is plated.
K	11		3.6.8 Control plates are used to check air quality and the sterility of the agar and the diluent.
K	9,21		3.6.9 Solidified plates are incubated at 35 ± 0.5°C for 48 ± 3 hours inverted and stacked no more than four high.
K	9		3.6.10 Quebec Colony Counter or its equivalent is used to provide the necessary magnification and visibility for counting plates.
K	1		3.6.11 A hand tally or its equivalent is used for accuracy in counting.
			3.7 Computation of Results -SPC
K	9		3.7.1 Colony counts determined in accordance with Part III, A, Sections 4.31 through 4.33 in <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , Fourth Edition.
C	19		3.7.2 Colony counts are reported as CFU/g of sample.
			3.8 Bacteriological Analysis of Shellfish Using the ETCP
C	2,3		3.8.1 Prepared modified MacConkey agar is used on the day that it is made.
K	3		3.8.2 Double strength modified MacConkey agar is used.
C	3		3.8.3 Prepared double strength modified MacConkey agar is heated to boiling, removed from the heat, and boiled again. This agar is never autoclaved.
K	2, 3		3.8.4 Twice boiled, double strength modified MacConkey agar and is maintained in a tempering bath at 45 to 50°C until used.
K	2, 3		3.8.5 Phosphate buffered saline is used as the sample diluent in the ETCP.
C	2, 3		3.8.6 The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
C	9		3.8.7 The sample homogenate is cultured within 2 minutes of blending.
C	2,3		3.8.8 Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed into a sterile container and the contents brought up to 60 mL with sterile, tempered phosphate buffered saline.
K	3		3.8.9 Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey Agar is added.
K	2,3, 22		3.8.10 The container is gently swirled or slowly inverted once to mix the contents, which are subsequently distributed uniformly over six plates.
C	1		3.8.11 Media and diluent sterility are determined with each use. Results are recorded and the records maintained.

€	1		3.8.12 Media productivity is determined using media appropriate properly diluted pour plated positive and negative control cultures for each batch of Modified MacConkey agar prepared. Positive control culture Negative control culture
C	3, 13		3.8.13 2 When solidified, the plates are placed inverted into an air incubator at $45.5 \pm 0.5^\circ\text{C}$ for 18 to 30 hours of incubation.
C	2		3.8.14 3 Plates are stacked no more than three high in the incubator.
C	2		3.8.15 4 Appropriately diluted pour plated process control cultures <i>accompany each set of samples throughout incubation.</i> The results are recorded and the records maintained. Positive process control _____ Negative process control _____
3.9 Computation of Results - ETCP			
K	11		3.9.1 Quebec Colony counter or its equivalent is used to provide the necessary magnification and visibility for counting.
O	1		3.9.2 A hand tally or its equivalent is used to aid in counting.
C	3, 6		3.9.3 All brick red colonies greater than 0.5 mm in diameter are totaled over all the plates and multiplied by a factor of 16.7.
C	3		3.9.4 Results are reported as CFU/100 grams of sample.
Bacteriological Examination of Soft-shelled Clams and American Oysters for Male Specific Coliphage (MSC)			
3.10 MSC Equipment and Supplies			
K	30		3.10.1 Sample containers used for the shucked sample are sterile, made of glass or some other inert material (i.e. polypropylene) and hold 100 – 125 mL.
C	27, 28		3.10.2 The refrigerated centrifuge used must have the capacity to accommodate the amount of shellfish sample required for the procedure, perform at 9000 x g and maintain a temperature of 4°C.
K	9		3.10.3 The level of water in the tempering bath covers the level of liquid and agar in the container or culture tubes.
C	27, 28		3.10.4 Sterile 0.22µm pore size syringe filters and pre-sterilized plastic or sterile glass syringes are used to sterilize the antibiotic solutions.
K	1		3.10.5 The sterility of each lot of pre-sterilized syringes and syringe filters is determined. Results are recorded and records maintained.
K	1		3.10.6 The sterility of each batch of reusable glass syringes is determined. Results are recorded and records maintained.
C	27, 28		3.10.7 The balance used provides a sensitivity of at least mg (0.01g.).
C	27, 28		3.10.8 The temperature of the incubator used is maintained at $36 \pm 1^\circ\text{C}$.
C	28		3.10.9 Sterile disposable 50 mL centrifuge tubes are used and their sterility is determined with each lot. Results are recorded and records maintained.
3.11 MSC Media Preparation			
K	28		3.11.1 Media preparation and sterilization is according to the validated method.
K	27, 28		3.11.2 Bottom agar, double strength soft agar and growth broth are prepared from their individual components.
K	27, 28		3.11.3 Soft agar is prepared double strength in volumes of 2.5 mL.
C	27, 28		3.11.4 The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
O	27, 28		3.11.5 Storage of the bottom agar under refrigeration does not exceed 1 month.
K	27, 28		3.11.6 Unsterilized soft agar is stored at -20°C - 15°C for up to 3 months.
K	27, 28		3.11.7 The soft agar is removed from the freezer and sterilized for 15 minutes at 121°C before use.
K	27, 28		3.11.8 Storage of growth broth in the refrigerator in loosely capped tubes/bottles does not exceed 1 month and in screw capped tubes/bottles does not exceed 3 months.
K	27, 28		3.11.9 Bottom agar plates are allowed to reach room temperature before use.

3.12 Preparation of the Soft-Shelled Clams and American Oysters for MSC Analysis			
K	2,11		3.12.1 Shucking knives, scrub brushes and blender jars are autoclave sterilized for 15 minutes prior to use.
O	2		3.12.2 The blades of shucking knives are not corroded.
O	9		3.12.3 The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
O	2		3.12.4 The faucet used for rinsing the shellfish does not contain an aerator.
K	9		3.12.5 The shellfish are scrubbed with a stiff, sterile brush and rinsed under tap water of drinking water quality.
O	9		3.12.6 The shellfish are allowed to drain in a clean container or on clean towels unlayered prior to shucking.
K	9		3.12.7 Immediately prior to shucking, the hands of the analyst are thoroughly washed with soap and water and rinsed in 70% alcohol or clean gloves are donned.
C	9		3.12.8 Shellfish are not shucked through the hinge.
C	9		3.12.9 The contents of shellfish (liquor and meat) are shucked into a sterile, tared blender jar or other sterile container.
K	9		3.12.10 A representative sample of at least 12 shellfish is used for the analysis.
K	2, 19		3.12.11 The sample is weighed to the nearest 0.1 gram.
3.13 MSC Sample Analysis			
C	28		3.13.1 <i>E. coli</i> Famp ATCC 700891 is the bacterial host strain used in this procedure.
K	27, 28		3.13.2 Host cell growth broth is tempered at $36 \pm 1^\circ\text{C}$ and vortexed (or shaken) to aerate prior to inoculation with host cells.
K	27, 28		3.13.3 Several host cell colonies are transferred to a tube of tempered, aerated growth broth and incubated at $36 \pm 1^\circ\text{C}$ for 4-6 hours to provide host cells in log phase growth for sample analysis.
C	27, 28		3.13.4 After inoculation, the host cell growth broth culture is not shaken.
C	28		3.13.5 A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.
C	28		3.13.6 The elution mixture is prepared w/v by weighing the sample and adding two equal portions of sterile growth broth by volume to the shellfish tissue.
C	28		3.13.7 The elution mixture is homogenized at high speed for 180 seconds.
C	28		3.13.8 Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
C	28		3.13.9 The homogenized elution mixture is centrifuged for 15 minutes at $9000 \times g$ at 4°C.
C	27, 28		3.13.10 The supernatant is pipetted off, weighed and the weight recorded.
C	27, 28		3.13.11 The supernatant is allowed to warm to room temperature about 20 to 30 minutes.
K	27, 28		3.13.12 The autoclaved soft agar is tempered and held at $51 \pm 1^\circ\text{C}$ throughout the period of sample analysis.
K	27, 28		3.13.13 Two hundred microliters (0.2 mL) of log phase host strain <i>E. coli</i> is added to the tempering soft agar immediately prior to adding the sample supernatant.
K	27, 28		3.13.14 The sample supernatant is shaken or vortexed before being added to the tempering soft agar.
C	27, 28		3.13.15 2.5 mL of sample supernatant is added to each tube of tempering soft agar.
C	27, 28		3.13.16 The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
C	27, 28		3.13.17 The soft agar/sample supernatant/host cell mixture is overlaid onto bottom agar plates and swirled gently to distribute the mixture evenly over the plate.
C	28		3.13.18 Ten (10) plates are used, 2.5 mL per plate for a total of 25 mL of supernatant analyzed per sample.

K	27, 28		3.13.19 Negative and positive control plates are prepared and accompany each set of samples analyzed. The results are recorded and records maintained. Positive control __
K	27, 28		3.13.20 Growth broth is used as the negative control or blank.
K	27, 28		3.13.21 Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately diluted to provide countable low levels of phage is used as the positive control.
K	2		3.13.22 A negative control plate is plated at the beginning and end of each set of samples analyzed.
K	27, 28		3.13.23 The positive control is plated after all the samples are inoculated and immediately prior to the final negative control.
C	27, 28		3.13.24 All plates are incubated at 36 ± 1°C for 18 ± 2 hours.
			3.14 Computation of Results - MSC
C	27		3.14.1 Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
C	28, 32		3.14.2 The working range of the method is 1 to 200 PFU per plate. When there are no plaques on all ten plates, the count is <6 PFU/100 grams for soft-shelled clams, <7 PFU/ 100 grams for American oysters, and <5 PFU/ 100 grams for quahog (hard) clams. If the density exceeds 200 PFU per plate on all plates, the count is given as > 20,000 PFU/100 grams.
K	28		3.14.3 The formula used for determining the density of MSC in PFU/100 grams is: (0.364) (N) (Ws), where N = total number of plaques counted on all 10 plates and Ws = weight of the supernatant used.
O	9		3.14.4 The MSC count is rounded off conventionally to give a whole number.

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PUBLIC HEALTH SERVICE U.S. FOOD AND DRUG ADMINISTRATION OFFICE OF FOOD SAFETY SHELLFISH AND AQUACULTURE POLICY BRANCH 5001 CAMPUS DRIVE COLLEGE PARK, MD 20740-3835 TEL. 240-402-2151/2055/4960 FAX 301-436-2601		
SHELLFISH LABORATORY EVALUATION CHECKLIST MARBIONC Brevetoxin (Neurotoxic Shellfish Poisoning; NSP) ELISA		
LABORATORY:		
ADDRESS:		
TELEPHONE:	FAX:	EMAIL:
DATE OF EVALUATION:	DATE OF REPORT:	LAST EVALUATION:
LABORATORY REPRESENTED BY:		TITLE:
LABORATORY EVALUATION OFFICER:		SHELLFISH SPECIALIST:
		REGION:
OTHER OFFICIALS PRESENT:		TITLE:
Items which do not conform are noted by:		
C – Critical K - Key O - Other NA - Not Applicable Conformity is noted by a “√”		

PART I – QUALITY ASSURANCE		
Code	REF	Item Description
1.1 Quality Assurance (QA) Plan		
K	3, 6	1.1.1 Written Plan adequately covers all the following: (check \checkmark those that apply) a. Organization of the laboratory. b. Staff training requirements. c. Standard operating procedures. d. Internal quality control measures for equipment, their calibration maintenance, repair, performance and rejection criteria established. e. Laboratory safety. f. Internal performance assessment. g. External performance assessment.
C	3	1.1.2 QA Plan is implemented.
1.2 Educational/Experience Requirements		
C	State's Human Resources Department	1.2.1 In state/county laboratories, the supervisor meets the state/county educational and experience requirements for managing a public health laboratory.
K	State's Human Resources Department	1.2.2 In state/county laboratories, the analyst(s) meets the state/county educational and experience requirements for processing samples in a public health laboratory.
C	USDA Microbiology & EELAP	1.2.3 In commercial/private laboratories, the supervisor must have at least a bachelor's degree or equivalent in microbiology, biology, chemistry, or another appropriate discipline with at least two years of laboratory experience.
K	USDA Microbiology & EELAP	1.2.4 In commercial/private laboratories, the analyst must have at least a high school diploma and shall have at least three months of experience in laboratory sciences.
1.3 Work Area		
O	3, 6	1.3.1 Adequate for workload and storage.
O	6	1.3.2 Clean and well lighted.
O	6	1.3.3 Adequate temperature control.
O	6	1.3.4 All work surfaces are nonporous and easily cleaned.
1.4 Laboratory Equipment		
O	4	1.4.1 The pH meter has a standard accuracy of 0.1 unit.
K	4	1.4.2 pH paper in the appropriate range (i.e., 1-4), if used, is used with minimum accuracy of 0.5 pH units.
K	3	1.4.3 The pH meter is calibrated daily when in use. Results are recorded, and records are maintained.
K	6	1.4.4 Effect of temperature has been compensated for by an ATC probe, use of a triode or by manual adjustment.
K	6	1.4.5 The pH meter manufacturer instructions are followed for calibration or a minimum of two standard buffer solutions (pH 7 and 10) is used to calibrate the pH meter. Standard buffer solutions are used once and discarded.
K	3, 7	1.4.6 Electrode acceptability is determined daily or with each use following either slope or millivolt procedure.
K	2, 4	1.4.7 The balances being used provide an appropriate sensitivity at the weights of use, at least 0.1 g for laboratory precision balances and 0.1 mg for analytical balances.
K	6	1.4.8 The balance calibration is checked monthly using NIST class S, ASTM class 1 or 2 weights or equivalent. Results are recorded, and records are maintained.
K	1	1.4.9 Refrigerator temperature is maintained between 0 and 4 °C.
K	6	1.4.10 Refrigerator temperature is monitored at least once daily <u>on workdays</u> . Results are recorded and records maintained.
K	11	1.4.11 Freezer temperature is maintained at -10 °C or below.
K	6	1.4.12 Freezer temperature is monitored at least once daily <u>on workdays</u> . Results are recorded and records maintained.

C	9		1.4.13 All in-service thermometers are properly calibrated and immersed.
K	5		1.4.14 All glassware is clean.
C	11		1.4.15 Absorbance Microplate reader equipped with filter for measurement at 450 nm is used.
O	2		1.4.16 Absorbance Microplate reader performance is evaluated at least annually using manufacturer instructions or a check standard microplate at the appropriate wavelength (450) to assess alignment, accuracy, reproducibility, and linearity. Method used:
K	2		1.4.17 Autopipettors are calibrated for the appropriate volumes used and checked annually for accuracy. Results are recorded, and records are maintained.
O	11		1.4.18 A centrifuge capable of holding 15 mL or 50 mL polypropylene tubes is used.
			1.5 Reagents and Reference Solution Preparation and Storage
C	11		1.5.1 All solvents and reagents used are ACS grade materials or better.
O	6		1.5.2 Water contains < 100 CFU/ml as determined monthly using the heterotrophic plate count method. Results are recorded, and records are maintained. (Not required for bottled reagent grade or HPLC grade water when used immediately upon opening. If the bottle of water is not used entirely immediately, the water must be tested as above prior to continued use.)
K	6		1.5.3 Reagents are properly stored and labeled with the date of receipt, date opened or date prepared and expiration date.
C	11		1.5.4 Brevetoxin-3 (BTX-3 or PbTx-3) provided with the MARBIONC ELISA kit is used as the reference standard.
C	11		1.5.5 Stock standard solution is made by diluting brevetoxin-3 reference standard to 1 µg/ml in 100% methanol in a volumetric flask.
C	11		1.5.6 Working standard solution (100 ng/ml) is made by diluting 1 ml of stock solution to 10 ml in a volumetric flask using 100% methanol.
K	11		1.5.7 Extraction solvent (80% methanol) is made by adding 800 ml of methanol to a 1 L graduated cylinder and bringing the total volume to 1 L with water.
K	11		1.5.8 Phosphate Buffered Saline, pH 7.4 and Phosphate Buffered Saline, 0.05% Tween 20, pH 7.4 are used within 1 week of preparation. pH of prepared media is determined to ensure it is consistent with manufacturers requirements. Results are recorded, and records are maintained.
K	11		1.5.9 Phosphate Buffered Saline, pH 7.4 and Phosphate Buffered Saline, 0.05% Tween 20, pH 7.4 are stored in refrigerator for no longer than 1 week and brought to room temperature before use.
K	11		1.5.10 Gelatin stock solution is prepared by dissolving 5 g gelatin in 100 ml water and stirring the solution over gentle heat on a stir plate until clear. Gelatin stock solution is aliquoted into smaller volumes (e.g., 15 ml centrifuge tubes) and refrigerated.
K	11		1.5.11 Blocking buffer is prepared by dissolving 1 pouch in 200 ml water. Blocking buffer solution is aliquoted into 50-ml centrifuge tubes and refrigerated.
K	11		1.5.12 PGT (PBS, 0.05% Tween, 0.5% gelatin) is made fresh daily by measuring 5 ml liquified gelatin stock solution into a 50-ml centrifuge tube and filling to 50 ml with PBS-Tween.
C	11		1.5.13 Stock and working standard solutions are stored -10 °C or below.
C	5		1.5.14 All standards used are within expiration date (or 1 year <u>from open date</u> if not provided).
			1.6 Collection and Transportation of Samples
O	4, 1		1.6.1 Shellstock are collected in clean, waterproof, puncture resistant containers.
K	4, 1		1.6.2 Samples are appropriately labeled with the collector's name, type of shellstock, the harvest area, and time and date of collection.

C	4, 1		1.6.3 Immediately after collection, shellstock samples are placed in dry storage (ice chest or equivalent) which is maintained between 0 and 10 °C with ice or cold packs for transport to the laboratory.
K	2, 10		1.6.4 Time from collection to initiation of the extraction should not exceed 24 hours. However, if significant delays are anticipated or if they occur, the laboratory has an appropriate contingency plan in place to handle the samples. For samples shipped live in accordance with 1.6.3, the contingency plan ensures samples remain within allowable temperature tolerances and animals are alive upon receipt. The contingency plan also addresses field and/or laboratory processing that ensures the integrity of the sample or extract until initiation of the assay. For example, samples are washed, shucked, drained and processed as follows: a. refrigerated or frozen until extracted; b. homogenized and frozen until extracted; or c. extracted, the supernatant decanted, and refrigerated or frozen until assayed.
C	2		1.6.5 Frozen shucked product or homogenates are allowed to thaw completely, and all liquid is included as part of the sample before being processed further.
PART II – ASSAY OF SHELLFISH FOR NSP TOXINS			
2.1 Preparation of Sample			
C	4		2.1.1 At least 12 animals are used per sample, or the laboratory has an appropriate contingency plan for dealing with non-typical species of shellfish.
O	4		2.1.2 The outside of the shell is thoroughly cleaned with fresh water.
O	4		2.1.3 Shellstock are opened by cutting the adductor muscles.
O	4		2.1.4 The inside surfaces of the shells are rinsed with fresh water to remove sand and other foreign materials.
O	4		2.1.5 Shellfish meats are removed from the shell by separating the adductor muscles and tissue connecting at the hinge.
C	4		2.1.6 Damage to the body of the mollusk is minimized in the process of opening.
O	4		2.1.7 Shucked shellfish are drained on a #10 mesh sieve or equivalent without layering for 5 minutes.
K	4		2.1.8 Pieces of shell and drainage are discarded.
C	2, 4		2.1.9 Drained meats or previously cooled/refrigerated shucked meats and their drip loss liquid or thawed homogenates with their freeze-thaw liquid are blended at high speed until homogenous (60-120 seconds).
2.2 Sample Extraction			
K	4		2.2.1 Sample homogenates are extracted as soon as possible (preferably the same day) or stored in the freezer.
C	11		2.2.2 One (1) gram of homogenized sample is weighed into a 15 ml or 50 ml polypropylene centrifuge tube and subsequently extracted.
C	11		2.2.3 The sample homogenate is extracted by adding 9 ml extraction solvent (80% aqueous methanol) and vortexing at highest speed for 2 minutes.
C	11		2.2.4 The homogenate/methanol mixture is centrifuged at a minimum of 3,000xg for 10 minutes.
C	11		2.2.5 The supernatant is transferred to a clean, labeled graduated 15-ml centrifuge tube and brought to a final volume of 10 ml with extraction solvent.
K	11		2.2.6 Crude extracts are sealed tightly in glass vials and stored at -10 °C or below until analyzed.

		2.3 Analysis
C	11	2.3.1 Only high-binding flat-bottom plates no older than 1 year are used (e.g., Nunc Maxisorp Immunoplates).
C	2, 11	2.3.2 A standard calibration curve of seven concentrations (0.078-5 ng PbTx-3/ml) is included on each plate. Results are recorded, and records are maintained.
C	11	2.3.3 2 When pipetting kit reagents that are pre-diluted in glycerol (A, C, and D): a. the pipet tip is not pre-rinsed, b. only the very tip of the pipet tip is inserted into the vial to withdraw the required amount, c. the tip is submerged into the buffer when dispensing and rinsed several times with buffer to ensure complete transfer
K	2	2.3.4 3 Crude The working standard solution and crude sample extracts are brought to room temperature and thoroughly mixed before withdrawing an aliquot for analysis use. TMB is brought to room temperature in the dark prior to Assay Step 7.
€	11	2.3.5 Crude extracts are diluted with PGT before analysis. The minimum dilution for shellfish extracts is 1:40 (25 µl + 975 µl PGT) (resulting in a sample dilution of 1:400).
€	11	2.3.6 For quantitative (actionable) results, serial dilutions (n=7) of each sample extract are assayed. Fewer dilutions are permissible for screening purposes only.
C	11	2.3.7 4 Assay Step 1: Reagent A is diluted by 300 (or as specified in kit instructions) in PBS, 100 µl is added to each well of the 96-well plate, and the plate is incubated on a plate shaker for 1 hour. After 1 hour, the liquid is poured from the plate, and all wells are rinsed 3 times with 300 µl PBS (no Tween for this wash step).
C	11	2.3.8 5 Assay Step 2: Each well is filled with 250 µl of blocking buffer. The plate is incubated on a plate shaker for 30 minutes. After 1 hour 30 minutes, the liquid is poured from the plate, and all wells are rinsed 3 times with 300 µl PBS-Tween.
C	11	2.3.9 6 Assay Step 3a: Crude sample extracts are diluted with PGT for analysis. The minimum dilution for shellfish extracts is 1:40 (25 µl + 975 µl PGT) (resulting in a sample dilution of 1:400). Serial dilutions (n=7) of each crude sample extract are prepared in PGT. and a standard calibration curve of seven concentrations (0.078-5.0 ng PbTx-3/ml) are prepared in PGT.
C	11	2.3.7 Assay Step 3b: A standard calibration curve of seven concentrations (0.078-5.0 ng PbTx-3/ml) is prepared in PGT and is included on each plate.
C	11	2.3.10 8 Assay Step 4: 100 µl of each sample or standard dilution is loaded on to the microplate as well as two reference wells (containing PGT only) adjacent to each set of sample dilutions. Each dilution of standard or sample is added to duplicate wells. Plate layout identifying locations of samples and standards on the plate is documented.
C	11	2.3.11 9 Assay Step 5: Reagent C is diluted by 300 (or as specified in kit instructions) in PGT, 100 µl is added to each well of the 96-well plate (which contains samples or standards), and the plate is incubated on a plate shaker for 90 minutes. After 90 minutes, the liquid is poured from the plate, and all wells are rinsed 3 times with 300 µl PBS-Tween.
C	11	2.3.12 10 Assay Step 6: Reagent D is diluted by 800 (or as specified in kit instructions) in PGT, 100 µl is added to each well of the 96-well plate, and the plate is incubated on a plate shaker for 1 hour. After 1 hour, the liquid is poured from the plate, all wells are rinsed 3 times with 300 µl PBS-Tween, and one final time with 300 µl PBS only to ensure no Tween remains on the plate. TMB should be brought to room temperature in the dark.
C	11	2.3.13 11 Assay Step 7: Each well is filled with 100 µl of room temperature TMB (3,3',5,5'-Tetramethylbenzidine) and incubated until a blue color develops in the reference wells. The reaction is stopped by adding 100 µl of 0.5M sulfuric

			acid solution to each well, and the absorbance in the wells at 450 nm is measured in a microplate reader.
K	11		2.3. 14 <u>12</u> Plates are covered with microplate sealing film during <u>all</u> a-H incubation steps (Steps 1-6 above).

C	11	2.3. 15 13 Plates are protected from light by covering with aluminum foil during color development (Step 7 above).
K	11	2.3. 16 14 The timing of the final step should be standardized is reassessed with each new lot of kit reagents and each new lot of TMB to achieve maximum optical densities (at 450 nm) of 1.0 +/- 30% to ensure that reference well absorbance values fall into an appropriate range.
		2.4 Quality Control
C	11	2.4.1 Acceptance of assay (plate) results is dependent on meeting the following criteria: a. Absorbance of standard reference wells (Amax) must be ≥ 0.6 . b. CV of raw absorbance of duplicate wells for standard curve within the linear range of the assay (30-70% inhibition) must be $< 20\%$.
C	11	2.4.2 Acceptance of individual sample results is dependent on meeting the following criteria: a. CV of raw absorbance of duplicate wells for sample dilutions used for quantitation (30-70% inhibition) must be $< 20\%$. b. CV of calculated concentrations of different sample dilutions within the linear range of the assay (30-70% inhibition) must be $< 20\%$.
		2.5 Calculation of Sample Toxin Concentration
C	11	2.5.1 Absorbance values are converted to % color inhibition: a. $\% \text{ inhibition} = [1 - (\text{Avg of duplicate } A / A_{\text{max}})] \times 100\%$ where Amax is the average absorbance of the reference wells oriented below the sample or standard dilutions
C	11	2.5.2 Using the 4-parameter logistic (4PL) curve in an appropriate software program, a curve is fit to the positive control with ng toxin/ml on the x-axis (log scale) and % inhibition on the y-axis (linear scale).
C	11	2.5.3 The concentrations for sample dilutions falling within 30%-70% inhibition are interpolated from the standard curve.
C	11	2.5.4 Sample toxin concentration is calculated by multiplying the interpolated concentration by the sample dilution factor and dividing by 1000 to obtain PbTx-3 eq. results in ppm.
C	11	2.5.5 If more than one dilution of a sample falls within 30%-70%, the mean of the two calculated concentrations is used.
C	8	2.5.6 A result of ≤ 1.6 ppm in clams and ≤ 1.8 ppm in oysters is considered negative and can substitute for testing by an Approved Method for the purposes of controlled relaying, controlled harvest end-product testing, or to re-open a previously closed area. A positive result (> 1.6 ppm in hard clams and sunray venus clams and > 1.8 ppm in oysters) requires additional testing by an Approved Method to support management actions.
O	8	2.5.7 Laboratory reports to the Shellfish Management Authority detail sample date, location, species (matrix), date tested, analyst name, result of ELISA in ppm, and the actionable ELISA threshold for the species tested.

REFERENCES
1. American Public Health Association. 1984. <i>Compendium for the Microbiological Examination of foods</i> , 2 nd Edition. APHA. Washington D.C.
2. Good Laboratory Practice.
3. Association of Official Analytical Chemists (AOAC). 1991. <i>Quality Assurance Principles for Analytical Laboratories</i> . AOAC, Arlington, VA.
4. American Public Health Association. 1970. <i>Recommended Procedures for the Examination of Sea Water and Shellfish</i> , 4 th Edition. APHA, Washington, D.C.
5. Consult reference standard product literature.
6. APHA/WEF/AWWA. 1992. <i>Standard Methods for the Examination of Water and Wastewater</i> , 18 th Edition. APHA, Washington, D.C.
7. Consult pH electrode product literature.
8. U.S. Food and Drug Administration (FDA) and Interstate Shellfish Sanitation Conference (ISSC). 2017. <i>NSSP Guide for the Control of Molluscan Shellfish</i> . FDA/ISSC, Washington, D.C. and Columbia, S.C.
9. U.S. Department of Commerce. 1976. NBS Monograph 150. U.S. Department of Commerce, Washington, D.C.
10. <i>Compendium of Methods for the Microbiological Examination of Foods</i> , 3 rd Edition, pg. 901.
11. MARBIONC Enzyme-linked Immunosorbent Assay (ELISA) for the determination of Neurotoxic Shellfish Poisoning (NSP) toxins in molluscan shellfish. (ISSC proposal 17-107, supporting documents Appendix A)

LABORATORY STATUS	
LABORATORY	DATE
LABORATORY REPRESENTATIVE:	
NEUROTOXIC SHELLFISH POISON (NSP or Brevetoxin) COMPONENT: PARTS I AND II	
A. Results Total # of Critical (C) Nonconformities _____ Total # of Key (K) Nonconformities _____ Total # of Critical, Key, and Other (O) Nonconformities _____	_____ _____ _____
B. Criteria for Determining Laboratory Status of the brevetoxin (NSP) ELISA Component	
1. Conforms Status: The NSP component of this Laboratory is in conformity with NSSP requirements if all of the following apply. <ul style="list-style-type: none"> a. No Critical nonconformities. b. and <6 Key nonconformities. c. and <12 Total nonconformities. 	
2. Provisionally Conforms Status: The NSP component of this laboratory is determined to be provisionally conforming to NSSP requirements if all of the following apply. <ul style="list-style-type: none"> a. the number of critical nonconformities is ≥ 1 but < 4. b. and < 6 Key nonconformities. c. and < 12 Total nonconformities. 	
3. Does Not Conform Status: The NSP component of this laboratory is not in conformity with NSSP requirements when any of the following apply. <ul style="list-style-type: none"> a. The total # of Critical nonconformities is ≥ 4. b. or the total # of Key nonconformities is ≥ 6. c. or the total # of Critical, Key, or Other is ≥ 12. 	
C. Laboratory Status (<i>circle appropriate</i>) <div style="text-align: center;"> Does Not Conform – Provisionally Conforms – Conforms </div>	
Acknowledgement by Laboratory Director/Supervisor: All corrective Action will be implemented and verifying substantiating documentation received by the Laboratory Evaluation Officer on or before _____. Laboratory Signature: _____ Date: _____ LEO Signature: _____ Date: _____	

Example 4

Oysters from a restricted harvest area produced the following end-product deputed fecal coliform counts which have been arrayed in ascending order for ease of calculation. In this case, there are 30 samples (n=30) because all 10 lots were under conditional protocol and required 3 end product samples each.

Sample #	FC Count (MPN/100 grams)
1	3.9 (<4)
2	3.9 (<4)
3	3.9 (<4)
4	3.9 (<4)
5	3.9 (<4)
6	3.9 (<4)
7	3.9 (<4)
8	3.9 (<4)
9	3.9 (<4)
10	3.9 (<4)
11	3.9 (<4)
12	4
13	4
14	4
15	4
16	7
17	7
18	12
19	12
20	12
21	16
22	16
23	16
24	22
25	28
26	35
27	35
28	56
29	101 (>100)
30	101 (>100)

In this example, $n=30$. Applying the formula $(n+1)P/100$ gives the position of the 90th percentile in the arrayed data: $30+1=31$, $31*90=2790/100=27.9$. Thus the 90th percentile for a thirty (30) sample array is the 27.9th sample in the array. The value for the 27.9th sample in the array is interpolated by subtracting the value of sample #27 which is 35 from the value of sample #28 which is 56 to give 21. 21 is then multiplied by 0.9 to give 18.9 which is added to the value of sample #27 which is 35 to give 53.9. Rounding this value off to 54, the value for the ninetieth (90th) percentile becomes 54.

Example 5

Once again, oysters from a restricted harvest area produced the following end-product depurated fecal coliform counts which have been arrayed in ascending order for ease of calculation. But in this case, there are 24 samples ($n=24$) because 6 of the lots were under conditional protocol and required 3 end product samples each, and the other 4 lots, under routine verification, required just one end product sample each.

Sample #	FC Count (MPN/100 grams)
1	3.9 (<4)
2	3.9 (<4)
3	3.9 (<4)
4	3.9 (<4)
5	3.9 (<4)
6	3.9 (<4)
7	3.9 (<4)
8	3.9 (<4)
9	4
10	4
11	4
12	7
13	7
14	7
15	7
16	12
17	12
18	12
19	16
20	28
21	28
22	36
23	101 (>100)
24	101 (>100)

In this example, $n=24$. Applying the formula $(n+1)P/100$ gives the position of the 90th percentile in the arrayed data: $24+1=25$, $25*90=2250/100=22.5$. Thus the 90th percentile for a twenty four (24) sample

array is the 22.5th sample in the array. The value for the 22.5th sample in the array is interpolated by subtracting the value of sample #22 which is 36 from the value of sample #23 which is 101 to give 65. 65 is then multiplied by 0.5 to give 32.5 which is added to the value of sample #22 which is 36 to give 68.5. Rounding this value off to 69, the value for the ninetieth (90th) percentile becomes 69.

Source*	Type of seafood	Species (if specified)	Nutritional Composition			Comment	Nutritional Composition					
			Protein (g)	Total Lipid (g)	Carbohydrate (g)		Protein (g)	Total Lipid (g)	Carbohydrate (g)			
Oysters	FDC	Pacific oyster	<i>Crassostrea gigas</i>	9.5	2.3	5.0		All Oysters				
	FDC	Eastern oyster	<i>Crassostrea virginica</i>	5.7	1.7	2.7		Average	9.3	2.1	4.1	
	V&G 2017	Eastern oyster		14.0	4.9			SD	3.6	1.0	1.7	
	V&G 2017	Oyster, mixed		18.8	3.6			RSD	38.9	48.1	41.6	
	FAO	Cupped oysters, flesh, raw		8.9	1.8	5.3		Oysters excluding "mixed"				
	FAO	Pacific cupped oyster, flesh, raw (n.s.)		9.7	1.8	5.8		Avg	8.6	1.7	4.1	
	FAO	Pacific cupped oyster, farmed, flesh, raw		9.6	1.6	7.3		SD	2.5	0.3	1.7	
	FAO	Mangrove cupped oyster, flesh, raw		8.4	1.7	3.3		RSD	29.1	16.4	41.0	
	FAO	American cupped oyster, flesh, raw (n.s.)		5.6	1.4	3.8						
	FAO	American cupped oyster, flesh, farmed, raw (USA)		5.1	1.4	4.7						
	FAO	American cupped oyster, flesh, wild, raw (USA)		6.0	1.2	3.4						
	FAO	Flat oysters, flesh, raw		9.4	2.1	2.6						
	FAO	European flat oyster, flesh, raw		10.8	1.6	0.9						
	Clams	FDC	Hard clam/southern quahog	<i>Mercenaria campechiensis</i>	5.7		2.9		All Clams (incl. Geoduck)			
		V&G 2017	Clam, mixed		25.5	1.9			Average	13.2	1.6	3.9
FAO		Venus clams, flesh, raw		9.2	1.2	3.5		SD	6.4	0.6	1.6	
FAO		Striped venus, flesh, raw		9.2	1.6	3.8		RSD	48.4	40.0	42.3	
							Clams excluding "mixed" and Geoduck					
							Avg	8.0	1.4	3.4		
Geoduck	Oliveira et al. 2011	Pacific geoduck - mantle	<i>Panopea abrupta</i>	14.3	2.6	6.9	%wt per wt	Geoduck				
	Oliveira et al. 2011	Pacific geoduck - siphon	<i>Panopea abrupta</i>	15.3	0.7	2.2	%wt per wt	Average	14.8	1.6	4.6	
								SD	0.5	0.9	2.4	
								RSD	3.4	57.2	51.8	
Scallops	FDC	Sea scallop (Chilean scallop)	<i>Argopecten purpuratus</i>	15.0	0.9	0.9		All Scallops				
	FDC	MAGDALENA BAY SCALLOPS	<i>Argopecten circularis</i>	15.0	0.9	6.2		Average	15.6	0.9	2.3	
	FDC	North Atlantic Sea Scallop	<i>Placopecten magellanicus?</i>	12.4	0.4	3.5		SD	2.8	0.2	1.8	
	FDC	Giant Sea Scallop	<i>Placopecten magellanicus</i>	15.0	0.9	0.9		RSD	17.9	25.9	78.1	
	FDC	Bay scallops		15.0	0.9	2.7						
	FDC	Sea scallop	<i>Placopecten magellanicus</i>	13.3	0.9	0.9						
	V&G 2017	Scallop, mixed		17.0	0.8							
	V&G 2017	Scallop, bay and sea		23.2	1.4							
	FAO	Scallops, flesh, raw		14.9	0.7	2.2						
	FAO	Great Atlantic scallop, flesh, raw		14.8	0.8	0.9						
Mussels	FDC	farmed Chilean mussel	<i>Mytilus chelensis</i>	13.9	2.5	4.1		All Mussels				
	FDC	Blue mussel	<i>Mytilus edulis</i>	11.9	2.2	3.7		Average	13.5	1.9	4.2	
	G&V 2009	estuarine mussels	<i>Perna viridis</i>	28.4	1.7	6.5	glycogen	SD	5.8	0.4	1.7	
	G&V 2009	estuarine mussels	<i>Meretrix meretrix</i>	24.5	1.5	3.9	glycogen	RSD	42.8	21.6	39.8	
	FAO	Mytilus mussels, flesh, raw		11.4	2.0	5.0		Mussels excluding estuarine				
	FAO	Korean mussel, flesh, raw	<i>Mytilus coruscus</i>	12.4	2.6	8.8		Avg	11.2	1.9	4.1	
	FAO	Blue mussel, flesh, raw	<i>Mytilus edulis</i>	11.8	1.9	3.9		SD	1.7	0.4	1.7	
	FAO	Mediterranean mussel, flesh, raw (n.s.)		9.6	1.8	3.4		RSD	15.1	21.4	41.5	
	FAO	Mediterranean mussel, wild, flesh, raw		10.2	1.8	3.5						
	FAO	Mediterranean mussel, farmed, flesh, raw		8.3	1.9	3.8						
	FAO	Perna mussel, flesh, raw		13.5	2.0	2.5						
	FAO	New Zealand mussel, flesh, raw (New Zealand)		10.7	1.8	3.9						
	FAO	Green mussel, flesh, raw		9.3	0.9	2.0						
	Cockle	FDC	cockle, Alaskan Native		13.5	0.7	4.7					

*Sources:

FDC USDA FoodDataCentral
FAO FAO/INFOODS Global Food Composition Database For Fish and Shellfish, version 1.0 (uFish1.0)
V&G 2017 V. Venugopal, K. Gopakumar 2017
G&V 2009 S. GOPALAKRISHNAN1 & K. VIJAYAVEL 2009
Oliveira et al. 2011 Oliveira et al. 2011

Compilation of State, County, and Local Anti-Idling Regulations

EPA420-B-06-004
April 2006

Compilation of State, County, and Local Anti-Idling Regulations

Transportation and Regional Programs Division
Office of Transportation and Air Quality
U.S. Environmental Protection Agency

The following compilation of state and local vehicle idling laws represents the U.S. Environmental Protection Agency's best efforts to catalogue, in one location, the variety of existing and proposed idling laws in their entirety. This document is for reference purposes only; please refer to the actual laws for requirements and compliance. This compilation may not include every state or local law, and you should enquire about your own jurisdiction's regulations on idling. We will make every effort to update this document when we are aware of new idling laws or changes to existing idling laws. For more information on state and local idling reduction laws, please visit the SmartWay Transport Partnership Web site at: www.epa.gov/smartway/idle-state.htm.

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Arizona

State Codes

ARIZONA REVISED STATUTES

§ 11-876. Engine idling restrictions; exemptions; applicability; civil penalty; definition

A. By July 1, 2002, a county that contains any portion of area A as defined in section 49-451 shall adopt, implement and enforce ordinances that place limits on the maximum idling time for engines that propel heavy-duty diesel vehicles with a gross vehicle weight rating of more than fourteen thousand pounds. The ordinances shall at least include exemptions for:

1. Certain types of vehicles, such as police, fire and other emergency vehicles.
2. Certain types of situations such as traffic delays or the need for a driver to sleep in the vehicle.
3. Certain types of equipment operations, such as refrigeration of cargo.

B. A county with a population of less than one million two hundred thousand persons shall adopt, implement and enforce the ordinances required by this section only for those portions of the county that are located in area A.

C. Any other county may adopt, implement and enforce ordinances that comply with this section.

D. A driver who violates an ordinance adopted pursuant to this section is subject to:

1. The imposition of a civil penalty of one hundred dollars for the first violation.
2. The imposition of a civil penalty of three hundred dollars for a second or any subsequent violation.

E. Ordinances adopted pursuant to this section may be enforced by a county control officer or any law enforcement officer who is authorized to enforce traffic laws. For violations of ordinances adopted pursuant to this section, an officer shall use a uniform civil ticket and complaint substantially similar to a uniform traffic ticket and complaint prescribed by the rules of procedure in civil traffic cases adopted by the supreme court. The officer may issue citations to persons who violate an ordinance adopted pursuant to this section.

F. In enforcing ordinances adopted pursuant to this section, a county control officer or authorized law enforcement officer shall only issue one citation per traffic stop or

investigation of a driver whose vehicle exceeds the maximum idling limits established pursuant to this section.

G. For the purposes of this section, "idling" means the operation of an engine in the operating mode where the engine is not engaged in gear, where the engine operates at a speed at the revolutions per minute specified by the engine or vehicle manufacturer for when the accelerator is fully released and there is no load on the engine.

Municipal Codes

MARICOPA COUNTY VEHICLE IDLING RESTRICTION ORDINANCE

SECTION 1 - GENERAL

A. **PURPOSE:** The Vehicle Idling Restriction Ordinance restricts, from idling for more than five (5) consecutive minutes, any device or combination of devices that meets all of the following criteria:

1. designed with a gross vehicle weight rating of more than 14,000 pounds; and
2. required under Arizona law (Arizona Revised Statute [ARS] Title 28 Chapters 7 and 9) to be registered; and
3. designed to operate on public highways; and
4. powered by a diesel engine.

B. **APPLICABILITY:** This Vehicle Idling Restriction Ordinance applies to vehicle idling within Maricopa County.

SECTION 2 - DEFINITIONS: For the purpose of this ordinance, the following definitions shall apply:

A. **COMBINATION OF DEVICES** – The coupling of two or more pieces of equipment that consist of the device which contains the diesel engine and an attached piece of equipment, which includes but is not limited to a trailer, cement mixer, refrigeration unit or automobile.

B. **DISTRIBUTION CENTER** – A place with multiple bays where vehicles load or unload materials.

C. **GROSS VEHICLE WEIGHT RATING** – The maximum vehicle weight for which the vehicle is designed as established by the manufacturer.¹

D. **IDLING** – The operation of a diesel engine when the engine is not engaged in gear.²

E. **POWER TAKE OFF (PTO) MECHANISM** – A unit that provides power from the engine to a trailer or other equipment.

¹ Mirrors the definition in R18-2-1001.36.

² Federal definition: "Curb-idle" means: (1) For manual transmission code light-duty trucks, the engine speed with the transmission in neutral or with the clutch disengaged. (2) For automatic transmission code light-duty trucks, curb-idle means the engine speed with the automatic transmission in the Park position (or Neutral position if there is no Park position); (2) For manual transmission code heavy-duty engines, the manufacturer's recommended engine speed with the clutch disengaged. For automatic transmission code heavy-duty engines, curb idle means the manufacturer's recommended engine speed with the automatic transmission in gear and the output shaft stalled.

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F. PRIMARY PROPULSION ENGINE – Any engine for which the primary function is to provide mechanical power to propel or direct a vehicle, regardless of whether that power is applied directly to the propeller shaft or indirectly by way of an electrical system.

G TRUCK STOP – A place of business that provides services to drivers and their vehicles in which the service time may exceed one (1) hour.

H. VEHICLE – Any device or combination of devices with a gross vehicle weight rating of more than 14,000 pounds, required under Arizona law (ARS Title 28 Chapters 7 and 9) to be registered, designed to operate on public highways and powered by a diesel engine.³

3 Note: AAC R18-2-101(69): "motor vehicle" means any self-propelled vehicle designed or transporting persons or property on public highways;

ARS 44-1301: "motor vehicle" means any automobile, motorcycle, truck, trailer, semitrailer, truck tractor and semitrailer combination or other vehicle operated on the roads of this state, used to transport person or property and propelled by power other than muscular power, but motor vehicle does not include traction engines, vehicles that run only on a track, bicycles or mopeds; ARS 49-541(16): "Vehicle" means any automobile, truck, truck tractor, motor bus or self-propelled or motor-driven vehicle registered or to be registered in this state and used upon the public highways of this state for the purpose of transporting persons or property, except implements of husbandry, road rollers or road machinery temporarily operated upon the highway.

ARS 49-581: "Motor vehicle" means any self-propelled vehicle including a car, van, bus or motorcycle and all other motorized vehicles;

ARS 28-101(29): "Motor vehicle": (a) means either: (i) A self-propelled vehicle; (ii) For the purposes of the laws relating to the imposition of a tax on motor vehicle fuel, a vehicle that is operated on the highways of this state and that is propelled by the use of motor vehicle fuel. (b) Does not include a motorized wheelchair or a motorized skateboard. For the purposes of this subdivision: (i) "motorized wheelchair" means a self-propelled wheelchair that is used by a person for mobility. (ii) "motorized skateboard" means a self-propelled device that has a motor, a deck on which a person may ride and at least two tandem wheel in contact with the ground.

ARS 28-101(50): "Truck" means a motor vehicle designed or used primarily for the carrying of property other than the effects of the driver or passengers and includes a motor vehicle to which has been added a box, a platform or other equipment for such carrying.

ARS 28-101 (51): "Truck tractor" means a motor vehicle that is designed and used primarily for drawing other vehicles and that is not constructed to carry a load other than a part of the weight of the vehicle and load drawn.

ARS 28-101 (52): "Vehicle" means a device in, on or by which a person or property is or may be transported or drawn on a public highway, excluding devices moved by human power or used exclusively on stationary rails or tracks.

ARS 28-101 (53): "Vehicle transporter" means either: (a) A truck tractor capable of carrying a load and drawing a semitrailer; (b) A truck tractor with a stinger-steered fifth wheel capable of carrying a load and drawing a semitrailer or a truck tractor with a dolly

mounted fifth wheel that is securely fastened to the truck tractor at two or more points and that is capable of carrying a load and drawing a semitrailer.

R17-4-435: "Motor carrier" as defined in ARS § 28-5201 except a motor carrier transporting passengers for hire in a vehicle with a design capacity of 6 or fewer persons. ARS 28-5201: "Motor vehicle" means a self-propelled motor driven vehicle or vehicle combination, except a lightweight motor vehicle, that is used on a public highway in the furtherance of a commercial enterprise. In research done by ADEQ, no definitions exist for "heavy duty motor vehicle," or "heavy duty diesel engine."

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SECTION 3 – REQUIREMENTS

A. **ORDINANCE** – No owner or operator of a vehicle shall permit the engine of such vehicle to idle for more than five (5) consecutive minutes except as provided in Section 4 (Exemptions) of this ordinance.

B. **VIOLATION** – Any owner or operator who violates this ordinance is subject to a civil penalty of \$100 for the first violation and \$300 for a second or any subsequent violation.⁴

C. **SIGN** – Each truck stop owner or operator and distribution center owner or operator shall erect and maintain a permanent sign(s) that is at least 12 inches by 18 inches in size indicating that the maximum idle time allowed in Maricopa County is 5 minutes. The sign(s) shall be posted in a conspicuous location, near the dispatcher, if applicable. In addition to the above, the sign shall at a minimum contain language outlining the following:

1. The County's vehicle idling information line, and
2. The amount of money the violator will be fined.

SECTION 4 – EXEMPTIONS: This ordinance shall not apply when:

- A. A vehicle is forced to remain motionless because of traffic or adverse weather conditions affecting the safe operation of the vehicle.
- B. A vehicle is being operated for emergency or law enforcement purposes.
- C. The primary propulsion engine of a vehicle meets all of the following criteria:
 1. is providing a power source necessary for mechanical operations other than propulsion; and
 2. involves a power take off (PTO) mechanism, or other mechanical device performing the same function as a PTO; and
 3. is powered by the engine for:
 - a. loading and unloading cargo, or
 - b. mixing or processing cargo, or
 - c. controlling cargo temperature, or
 - d. providing a mechanical extension to perform work functions.
- D. The primary propulsion engine of a vehicle is being operated at idle to conform to manufacturer’s warm up and cool down specifications, for maintenance or diagnostic purposes, or by manufacturers engaging the engines in testing for research and development.

⁴ Attorney General's Office (AGO) interpretation is that ARS Title 28 allows any law enforcement officer to enforce ARS 11-876, which authorized this ordinance, on private and/or public property.

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E. The primary propulsion engine of a vehicle is being operated to supply heat or air conditioning necessary for passenger comfort/safety in those vehicles operating for commercial passenger transportation or school purposes up to a maximum of 30 minutes/hour. If ambient temperatures exceed 75 degrees Fahrenheit, passenger buses are allowed to idle up to a maximum of 60 minutes in any 90-minute time period.

F. The primary propulsion engine of a vehicle is being operated to comply with the U.S. Code of Federal Regulation 49 CFR Part 395 and the Arizona Department Of Transportation (DOT) regulation R17-5-202 referencing hours of service restrictions.

California

State Codes

California Health & Safety Code

§ 40720. Operation in manner that does not cause engines on trucks to idle or queue for more than 30 minutes

(a) Each marine terminal in the state shall operate in a manner that does not cause trucks to idle or queue for more than 30 minutes while waiting to enter the gate into the marine terminal.

(1) Any owner or operator of a marine terminal that operates in violation of this subdivision is subject to a two hundred fifty dollar (\$ 250) fine per vehicle per violation.

(2) Marine terminals in the state shall be monitored by the district with jurisdiction over that terminal to ensure compliance with this subdivision.

(3) Citations for violations of this subdivision shall be issued by the applicable district, and shall include the truck license plate number or other unique identifier, which may include, but is not limited to, the cargo container number, the name of the marine terminal and port at which the violation occurred, and the date and time of the violation.

(4) Any action taken by the marine terminal to assess, or seek reimbursement from, the driver or owner of a truck for a violation of this subdivision shall constitute a violation of Article 3 (commencing with Section 42400) of Chapter 4 of Part 4.

(5) Any owner or operator of a marine terminal or port, or any agent thereof, who takes any action intended to avoid or circumvent the requirements of this subdivision or to avoid or circumvent the reduction of emissions of particulate matter from idling or queuing trucks is subject to a seven hundred fifty dollar (\$ 750) fine per vehicle per violation, including, but not limited to, either of the following actions:

(A) Diverting an idling or queuing truck to area freeways or alternate staging areas, including, but not limited to, requiring a truck to idle or queue inside the gate of a marine terminal.

(B) Requiring or directing a truckdriver to turn on and off an engine on a truck while that truck is idling or queuing.

(6) The owner or operator of a marine terminal does not violate this subdivision by causing a truck to idle or queue for more than 30 minutes while waiting to enter the gate into the marine terminal, if the delay is caused by acts of God, strikes, or declared state and federal emergencies, or if the district finds that an unavoidable or unforeseeable event caused a truck to idle or queue and that the terminal is in good faith compliance with this section.

(7) Failure to pay a fine imposed pursuant to paragraph (1) or (5) shall constitute a violation of Article 3 (commencing with Section 42400) of Chapter 4 of Part 4.

(b)(1) Subdivision (a) does not apply to any marine terminal that provides, as determined by the district, two continuous hours of uninterrupted, fully staffed receiving and delivery gates two hours prior to and after, peak commuter hours each day, at least five days per week.

(2) For the purposes of this subdivision, "peak commuter hours" shall be those hours determined by the district, in consultation with the owners and operators of the marine

terminals within the jurisdiction of each district and any labor union that is represented at those marine terminals. The district shall notify the marine terminals of the final determination of the peak commuter hours.

(c) Subdivision (a) does not apply to any marine terminal that operates fully staffed receiving and delivery gates for 65 hours, five days per week, if that marine terminal is located at a port that processes less than 3 million containers (20-foot equivalent units (TEUs)) annually.

(d) Subdivision (a) does not apply to any marine terminal that operates fully staffed receiving and delivery gates for 70 hours, five days per week, if that marine terminal is located at a port that processes more than 3 million containers (20-foot equivalent units (TEUs)) annually.

(e) The district shall determine the necessary level of monitoring and enforcement commensurate with the level of the truck idling or queuing problem existing within its jurisdiction.

(f) For the purposes of this section, "marine terminal" means a facility that meets all of the following criteria:

- (1) Is located at a bay or harbor.
- (2) Is primarily used for loading or unloading containerized cargo onto or off of a ship or marine vessel.
- (3) Contains one or more of the following:
 - (A) Piers.
 - (B) Wharves.
 - (C) Slips.
 - (D) Berths.
 - (E) Quays.
- (4) Is located at a port that processes 100,000 or more containers (20-foot equivalent units (TEUs)) annually.

(g) Notwithstanding paragraph (1) of subdivision (a), if a marine terminal implements a scheduling or appointment system for trucks to enter the terminal, the terminal shall be subject to a fine pursuant to subdivision (a) only for a truck that makes use of the system and that idles or queues for more than 30 minutes while waiting to enter the gate into the terminal, commencing from the start of the appointment or the time the truck arrives, whichever is later. The scheduling or appointment system shall meet all of the following requirements:

- (1) Provide appointments on a first-come-first-served basis.
- (2) Provide appointments that last at least 60 minutes and are continuously staggered throughout the day.
- (3) Not discriminate against any motor carrier that conducts transactions at the marine terminal in scheduling appointments.
- (4) Not interfere with a double transaction once inside the gate.
- (5) Not turn away or fine a motor carrier if that motor carrier misses an appointment.

California Health & Safety Code

§ 41700. Prohibited discharges

Except as otherwise provided in Section 41705, no person shall discharge from any

source whatsoever such quantities of air contaminants or other material which cause injury, detriment, nuisance, or annoyance to any considerable number of persons or to the public, or which endanger the comfort, repose, health, or safety of any such persons or the public, or which cause, or have a natural tendency to cause, injury or damage to business or property.

§ 42403.5. Discharge from idling engine of diesel-powered bus

(a) Notwithstanding Section 42407, any violation of Section 41700 resulting from the engine of any diesel-powered bus while idling shall subject the owner to civil penalties assessed under this article, which may be recovered pursuant to Section 42403 by the Attorney General, by any district attorney, or by the attorney for any district in which the violation occurs in any court of competent jurisdiction.

(b) There is no liability under subdivision (a) if the person accused of the violation establishes by affirmative defense that the extent of the harm caused does not exceed the benefit accrued to bus passengers as a result of idling the engine.

California Code of Regulations

§ 2480. Airborne Toxic Control Measure to Limit School Bus Idling and Idling at Schools

(a) Purpose. This airborne toxic control measure seeks to reduce public exposure, especially school age children's exposure, to diesel exhaust particulate matter and other toxic air contaminants by limiting unnecessary idling of specified vehicular sources.

(b) Applicability. Except as provided in subsection (d), this section applies to the operation of every school bus, transit bus, school pupil activity bus, youth bus, general public paratransit vehicle, and other commercial motor vehicle as defined in subsection (h).

(c) Idling Control Measure.

(1) A driver of a school bus, school pupil activity bus, youth bus, or general public paratransit vehicle:

(A) must turn off the bus or vehicle engine upon stopping at a school or within 100 feet of a school, and must not turn the bus or vehicle engine on more than 30 seconds before beginning to depart from a school or from within 100 feet of a school; and

(B) must not cause or allow a bus or vehicle to idle at any location greater than 100 feet from a school for:

(i) more than five consecutive minutes; or

(ii) a period or periods aggregating more than five minutes in any one hour.

(2) A driver of a transit bus or of a commercial motor vehicle not identified in (c)(1):

(A) must turn off the bus or vehicle engine upon stopping at a school and must not turn the bus or vehicle engine on more than 30 seconds before beginning to depart from a school; and

(B) must not cause or allow a bus or vehicle to idle at any location within 100 feet of, but not at, a school for:

(i) more than five consecutive minutes; or

(ii) a period or periods aggregating more than five minutes in any one hour.

(3) A motor carrier of a school bus, school pupil activity bus, youth bus, or general public paratransit vehicle must ensure that:

(A) the bus or vehicle driver, upon employment and at least once per year thereafter, is informed of the requirements in (c)(1), and of the consequences, under this section and the motor carrier's terms of employment, of not complying with those requirements;

(B) all complaints of non-compliance with, and enforcement actions related to, the requirements of (c)(1) are reviewed and remedial action is taken as necessary; and

(C) records of (3)(A) and (B) are kept for at least three years and made available or accessible to enforcement personnel as defined in subsection (g) within three business days of their request.

(4) A motor carrier of a transit bus or of a commercial motor vehicle not identified in (c)(1) must ensure that:

(A) the bus or vehicle driver, upon employment and at least once per year thereafter, is informed of the requirements in (c)(2), and of the consequences, under this section and the motor carrier's terms of employment, of not complying with those requirements;

(B) all complaints of non-compliance with, and enforcement actions related to, the requirements of (c)(2) are reviewed and remedial action is taken as necessary; and

(C) records of (4)(A) and (B) are kept for at least three years and made available or accessible to enforcement personnel as defined in subsection (g) within three business days of their request.

(d) Exemptions

This section does not apply for the period or periods during which:

(1) idling is necessary while stopped:

(A) for an official traffic control device;

(B) for an official traffic control signal;

(C) for traffic conditions over which the driver has no control, including, but not limited to: stopped in a line of traffic; or

(D) at the direction of a peace officer;

(2) idling is necessary to ascertain that the school bus, transit bus, school pupil activity bus, youth bus, general public paratransit vehicle, or other commercial motor vehicle is in safe operating condition and equipped as required by all provisions of law, and all equipment is in good working order, either as part of the driver's daily vehicle inspection, or as otherwise needed;

(3) idling is necessary for testing, servicing, repairing, or diagnostic purposes;

(4) idling is necessary, for a period not to exceed three to five minutes (as per the recommendation of the manufacturer), to cool down a turbo-charged diesel engine before turning the engine off;

(5) idling is necessary to accomplish work for which the vehicle was designed, other than transporting passengers, for example:

(A) collection of solid waste or recyclable material by an entity authorized by contract, license, or permit by a school or local government;

(B) controlling cargo temperature; or

(C) operating a lift, crane, pump, drill, hoist, mixer, or other auxiliary equipment other than a heater or air conditioner;

(6) idling is necessary to operate:

(A) a lift or other piece of equipment designed to ensure safe loading, unloading, or transport of persons with one or more disabilities; or

(B) a heater or an air conditioner of a bus or vehicle that has, or will have, one or more children with exceptional needs aboard;

(7) idling is necessary to operate defrosters, heaters, air conditioners, or other equipment to ensure the safety or health of the driver or passengers, or as otherwise required by federal or State motor carrier safety regulations;; or

(8) idling is necessary solely to recharge a battery or other energy storage unit of a hybrid

electric bus or vehicle.

(e) Relationship to Other Law

Nothing in this section allows idling in excess of other applicable law, including, but not limited to:

- (1) Title 13 California Code of Regulations Section 1226;
- (2) Vehicle Code Section 22515; or
- (3) any local ordinance or requirement as stringent as, or more stringent than, this section.

(f) Penalties

(1) For each violation of subsection (c)(1), a driver of a school bus, school pupil activity bus, youth bus, or general public paratransit vehicle is subject to a minimum civil penalty of 100 dollars and to criminal penalties to the maximum extent provided by law.

(2) For each violation of subsection (c)(2), a driver of a transit bus or other commercial motor vehicle is subject to a minimum civil penalty of 100 dollars and to criminal penalties to the maximum extent provided by law.

(3) For each violation of subsection (c)(3), a motor carrier of a school bus, school pupil activity bus, youth bus, or general public paratransit vehicle is subject to a minimum civil penalty of 100 dollars and to criminal penalties to the maximum extent provided by law.

(4) For each violation of subsection (c)(4), a motor carrier of a transit bus or other commercial motor vehicle is subject to a minimum civil penalty of 100 dollars and to criminal penalties to the maximum extent provided by law.

(g) Enforcement. This section may be enforced by the Air Resources Board, peace officers as defined in California Penal Code, title 3, chapter 4.5, Sections 830 et seq. and their respective law enforcement agencies' authorized representatives, and air pollution control or air quality management districts.

(h) Definitions.

The following terms are defined for the purposes of this section:

(1) Children With Exceptional Needs. "Children with exceptional needs" means children meeting eligibility criteria described in Education Code Section 56026.

(2) Commercial Motor Vehicle. "Commercial Motor Vehicle" means any vehicle or combination of vehicles defined in Vehicle Code Section 15210(b) and any other motor truck with a gross vehicle weight rating of 10,001 pounds or more, with the following

exceptions:

(A) a zero emission vehicle; or

(B) a pickup truck defined in Vehicle Code Section 471.

(3) Driver. "Driver" means any person who drives or is in actual physical control of a vehicle.

(4) General Public Paratransit Vehicle. "General public paratransit vehicle" means any motor vehicle defined in Vehicle Code Section 336, other than a zero emission general public paratransit vehicle, that is transporting school pupils at or below the 12th grade level to or from public or private schools or public or private school activities.

(5) Gross Vehicle Weight Rating. "Gross vehicle weight rating" means the weight specified by the manufacturer as the loaded weight of a single vehicle.

(6) Hybrid Electric Bus or Vehicle. "Hybrid electric bus or vehicle" means any school bus, transit bus, school pupil activity bus, youth bus, general public paratransit vehicle, or other commercial motor vehicle equipped with at least the following two sources of motive energy on board:

(A) an electric drive motor that must be used to partially or fully drive the bus or vehicle wheels; and

(B) one of the following:

(i) an internal combustion engine;

(ii) a turbine; or

(iii) a fuel cell.

(7) Idling. "Idling" means the engine is running while the bus or vehicle is stationary.

(8) Motor Carrier. "Motor carrier" means the registered owner, lessee, licensee, school district superintendent, or bailee of any school bus, transit bus, school pupil activity bus, youth bus, general public paratransit vehicle, or other commercial motor vehicle who operates or directs the operation of any such bus or vehicle on either a for-hire or not-for-hire basis.

(9) Motor Truck. "Motor truck" or "motortruck" means a motor vehicle designed, used, or maintained primarily for the, transportation of property.

(10) Official Traffic Control Device. "Official traffic control device" means any sign, signal, marking or device, consistent with Section 21400 of the Vehicle Code, placed or

erected by authority of a public body or official having jurisdiction, for the purpose of regulating, warning, or guiding traffic, but does not include islands, curbs, traffic barriers, speed humps, speed bumps, or other roadway design features.

(11) Official Traffic Control Signal. "Official traffic control signal" means any device, whether manually, electrically, or mechanically operated, by which traffic is alternately directed to stop and proceed and which is erected by authority of a public body or official having jurisdiction.

(12) School. "School" means any public or private school used for the purposes of education and instruction of more than 12 school pupils at or below the 12th grade level, but does not include any private school in which education and instruction is primarily conducted in private homes. The term includes any building or structure, playground, athletic field, or other area of school property. The term excludes unimproved school property.

(13) School Bus. "School bus" means any school bus defined in Vehicle Code Section 545, except a zero emission school bus.

(14) School Pupil Activity Bus. "School pupil activity bus" means any bus defined in Section 546 of the Vehicle Code, except a zero emission school pupil activity bus.

(15) Transit Bus. "Transit bus" means any bus defined in Vehicle Code Section 642, except a zero emission transit bus.

(16) Youth Bus. "Youth bus" means any bus defined in Vehicle Code Section 680, except a zero emission youth bus.

(17) Zero Emission School Bus, Transit Bus, School Pupil Activity Bus, Youth Bus, General Public Paratransit Vehicle, or Other Commercial Motor Vehicle. A "zero emission school bus, transit bus, school pupil activity bus, youth bus, general public paratransit vehicle, or other commercial motor vehicle" means any bus or vehicle certified to zero-emission standards.

California Code of Regulations

§ 2485. Airborne Toxic Control Measure to Limit Diesel-Fueled Commercial Motor Vehicle Idling

(a) Purpose. The purpose of this airborne toxic control measure is to reduce public exposure to diesel particulate matter and other air contaminants by limiting the idling of diesel-fueled commercial motor vehicles.

(b) Applicability. This section applies to diesel-fueled commercial motor vehicles that operate in the State of California with gross vehicular weight ratings of greater than 10,000 pounds that are or must be licensed for operation on highways. This specifically

includes:

(1) California-based vehicles; and

(2) Non-California-based vehicles.

(c) Requirements.

On or after February 1, 2005, the driver of any vehicle subject to this section:

(1) shall not idle the vehicle's primary diesel engine for greater than 5.0 minutes at any location, except as noted in Subsection (d); and

(2) shall not operate a diesel-fueled auxiliary power system (APS) to power a heater, air conditioner, or any ancillary equipment on that vehicle during sleeping or resting in a sleeper berth for greater than 5.0 minutes at any location when within 100 feet of a restricted area, except as noted in Subsection (d).

(d) Exceptions.

Subsection (c) does not apply for the period or periods during which

(1) a bus is idling for

(A) up to 10.0 minutes prior to passenger boarding, or

(B) when passengers are onboard;

(2) idling of the primary diesel-engine is necessary to power a heater, air conditioner, or any ancillary equipment during sleeping or resting in a sleeper berth. This provision does not apply when operating within 100 feet of a restricted area;

(3) idling when the vehicle must remain motionless due to traffic conditions, an official traffic control device, or an official traffic control signal over which the driver has no control, or at the direction of a peace officer, or operating a diesel-fueled APS at the direction of a peace officer;

(4) idling when the vehicle is queuing that at all times is beyond 100 feet from any restricted area;

(5) idling of the primary engine or operating a diesel-fueled APS when forced to remain motionless due to immediate adverse weather conditions affecting the safe operation of the vehicle or due to mechanical difficulties over which the driver has no control;

(6) idling to verify that the vehicle is in safe operating condition as required by law and that all equipment is in good working order, either as part of a daily vehicle inspection or

as otherwise needed, provided that such engine idling is mandatory for such verification;

(7) idling of the primary engine or operating a diesel-fueled APS is mandatory for testing, servicing, repairing, or diagnostic purposes;

(8) idling when positioning or providing a power source for equipment or operations, other than transporting passengers or propulsion, which involve a power take off or equivalent mechanism and is powered by the primary engine for:

(A) controlling cargo temperature, operating a lift, crane, pump, drill, hoist, mixer (such as a ready mix concrete truck), or other auxiliary equipment;

(B) providing mechanical extension to perform work functions for which the vehicle was designed and where substitute alternate means to idling are not reasonably available; or

(C) collection of solid waste or recyclable material by an entity authorized by contract, license, or permit by a school or local government;

(9) idling of the primary engine or operating a diesel-fueled APS when operating defrosters, heaters, air conditioners, or other equipment solely to prevent a safety or health emergency;

(10) idling of the primary engine or operating a diesel-fueled APS by authorized emergency vehicles while in the course of providing services for which the vehicle is designed;

(11) idling of military tactical vehicles during periods of training; and

(12) idling when operating equipment such as a wheelchair or people assist lift as prescribed by the Americans with Disabilities Act;

(e) Relationship to Other Law.

Nothing in this section allows idling in violation of other applicable law, including, but not limited to:

(1) California Vehicle Code Section 22515;

(2) Title 13, Section 2480, California Code of Regulations;

(3) California Health and Safety Code Section 40720; or

(4) any applicable ordinance, rule, or requirement as stringent as, or more stringent than, this section.

(f) Enforcement. This section may be enforced by the Air Resources Board; peace

officers as defined in California Penal Code, title 3, chapter 4.5, Sections 830 et seq. and their respective law enforcement agencies' authorized representatives; and air pollution control or air quality management districts.

(g) Penalties. For violations of subsection (c)(1) or (c)(2), the driver of a subject vehicle is subject to a minimum civil penalty of 100 dollars and to criminal penalties as specified in the Health and Safety Code and the Vehicle Code.

(h) Definitions.

The following definitions apply to this section:

- (1) "Authorized emergency vehicle" is as defined in Vehicle Code Section 165.
- (2) "Auxiliary power system" or "APS" means any device that provides electrical, mechanical, or thermal energy to the primary diesel engine, truck cab, or sleeper berth, as an alternative to idling the primary diesel engine.
- (3) "Bus" means any vehicle defined in Title 13, California Code of Regulations, Section 2480, subsections (h) (13)-(16), inclusive or as defined in the Vehicle Code Section 233.
- (4) "Commercial Motor Vehicle" means any vehicle or combination of vehicles defined in Vehicle Code Section 15210(b) and any other motor truck or bus with a gross vehicle weight rating of 10,001 pounds or more, except the following:
 - (A) a zero emission vehicle; or
 - (B) a pickup truck as defined in Vehicle Code Section 471.
- (5) "Driver" is as defined in Vehicle Code Section 305.
- (6) "Gross vehicle weight rating" is as defined in Vehicle Code Section 350.
- (7) "Highway" is as defined in Vehicle Code Section 360.
- (8) "Idling" means the vehicle engine is running at any location while the vehicle is stationary.
- (9) "Motor truck" or "motortruck" means a motor vehicle designed, used, or maintained primarily for the transportation of property.
- (10) "Official traffic control device" is as defined in Vehicle Code Section 440.
- (11) "Official traffic control signal" is as defined in Vehicle Code Section 445.
- (12) "Owner" is as defined in Vehicle Code Section 460.

(13) "Primary diesel engine" means the diesel-fueled engine used for vehicle propulsion.

(14) "Queuing" means (A) through (C)

(A) the intermittent starting and stopping of a vehicle;

(B) while the driver, in the normal course of doing business, is waiting to perform work or a service; and

(C) when shutting the vehicle engine off would impede the progress of the queue and is not practicable.

(D) Queuing does not include the time a driver may wait motionless in line in anticipation of the start of a workday or opening of a location where work or a service will be performed.

(15) "Restricted area" means any real property zoned for individual or multifamily housing units that has one or more of such units on it.

(16) "Safety or health emergency" means:

(A) a sudden, urgent, or usually unforeseen, occurrence; or

(B) a foreseeable occurrence relative to a medical or physiological condition.

(17) "Sleeper berth" is as defined in Title 13, California Code of Regulations, Section 1265.

(18) "Vehicle" is as defined in the Vehicle Code Section 670.

Municipal Codes

Auburn Municipal Code

LIMITATION ON ENGINE IDLING

§ 71.75 FINDINGS AND PURPOSE.

The City Council finds that:

(A) Air pollution is a public health concern in California. The Sacramento Region is currently designated as non-attainment for the 1-hour federal ozone standard, as well as the more stringent state ozone standard. Air pollution can cause or aggravate long illnesses such as acute respiratory infections, asthma, chronic bronchitis, emphysema, and lung cancer. In addition to health impacts, air pollution imposes significant economic costs and negative impacts on our quality of life (nuisance).

(B) Exhaust from motor vehicles (both on- and off-road) is a substantial source of ozone precursors in the Sacramento Region. Vehicle exhaust is also a source of carbon monoxide, particulate matter, toxic air contaminants, and greenhouse gases. Although new engines have become cleaner due to improved emission control technologies, the slow turnover in their inventory and the number of miles/hours these vehicles idle each year is hindering progress in improving regional air quality.

(C) Public agencies can play an important role in improving air quality by limiting the amount of time engines are allowed to idle within their jurisdiction. Public agencies have the responsibility to lead the effort to improve air quality by adopting ordinances that are cost-effective in reducing ozone precursor emissions and toxic air contaminants. This subchapter is based on and derived from the Sacramento Ozone Summit Model Green Contracting Ordinance.

(D) A study of idling exhaust emissions conducted by the U.S. Environmental Protection Agency (EPA420-R-02-025, October 2002) indicates that a typical 1980's-2001 model year truck operating on diesel fuel emits 144 grams per hour of nitrogen oxide and 8,224 grams per hour of carbon dioxide emissions and consumes about 0.82 gallons of diesel while idling.

(E) TIAX, a consultant for the Sacramento Metropolitan Air Quality Management District, estimated idling exhaust emissions from Heavy Duty diesel trucks (HHDV), Medium Heavy Duty Diesel Trucks (MHDV) and off-road construction equipment to be 2.3 tons per day of nitrogen oxide emissions and .23 tons per day of reactive organic gas emissions (Control Measure OFMS 52 & ONMS 45, April 2003). The maximum emissions reductions from full implementation of the Limitation on Engine idling Ordinance in the Sacramento Region was estimated to be 1.725 tons per day of nitrogen oxides emissions and .173 tons per day of reactive organic gas emissions (assuming a 75% compliance).

(F) Under this subchapter, a limitation on engine idling is established by the City of Auburn to discourage the idling of engines in the city.

(Ord. 04-5, eff. 8-10-2004)

§ 71.76 DEFINITIONS.

For the purpose of this subchapter, the following definitions shall apply unless the context clearly indicates or requires a different meaning.

COMMERCIAL MOTOR VEHICLE. Any on- road motor vehicle with a manufacturer's gross vehicle weight rating greater than 26,000 pounds, or as defined in Cal. Motor Vehicle Code § 15210(b).

DRIVER. Any person who drives, operates, or is in actual physical control of a vehicle.

EMERGENCY. A sudden, urgent, usually unforeseen occurrence.

EQUIPMENT OPERATOR. Any person who is in actual physical control of a piece of off-road equipment.

GROSS VEHICLE WEIGHT RATING. The weight specified by the manufacturer as the loaded weight of a single vehicle.

IDLING. The engine is running while the vehicle is stationary or the piece of off-road equipment is not performing work.

MEDIUM DUTY VEHICLE. Any motor vehicle with a manufacturer's gross vehicle weight rating of 6,001-14,000 pounds.

OFFICIAL TRAFFIC CONTROL DEVICE. Any sign, signal, marking or device, consistent with Cal. Vehicle Code § 21400, placed or erected by authority of a public body or official having jurisdiction, for the purpose of regulating, warning, or guiding traffic, but does not include islands, curbs, traffic barriers, speed humps, speed bumps or other roadway design features.

OFFICIAL TRAFFIC CONTROL SIGNAL. Any device, whether manually, electrically, or mechanically operated, by which traffic is alternately directed to stop and proceed and which is erected by authority of a public body or official having jurisdiction.

OFF-ROAD DIESEL EQUIPMENT. All non- road equipment with a horsepower rating of 70 or greater.

TRANSPORT REFRIGERATION UNIT or TRU. A refrigeration system powered by an engine designed to control the environment of temperature sensitive cargo. A TRU is a piece of off-road equipment regardless of its horsepower rating.

VEHICLE. Any on-road, self-propelled vehicle that is required to be registered and have a license plate by the Department of Motor Vehicles.

VEHICLE/EQUIPMENT OWNER. The registered owner, lessee, licensee or bailee of any heavy- or medium-duty vehicle or piece of off-road equipment who operates or directs the operation of any such vehicle or equipment on either a for hire or not for hire basis.

(Ord. 04-5, eff. 8-10-2004)

§ 71.77 APPLICABILITY.

This subchapter applies to the operation of all diesel fueled commercial vehicles over 26,000 lbs. Gross vehicle weight rating, and all off-road diesel powered equipment over

70 horsepower rating, except as provided in § 71.79. Additionally, this subchapter applies to TRU engines as specified in § 71.76.

(Ord. 04-5, eff. 8-10-2004)

§ 71.78 IDLING.

(A) A driver of a vehicle:

- (1) Must turn off the engine upon stopping at a destination; and
- (2) Must not cause or allow an engine to idle at any location for more than five consecutive minutes.

(B) An equipment operator of an off-road piece of equipment not identified in subdivision (A)(1) above must not cause or allow an off-road piece of equipment to idle at any location for more than five consecutive minutes.

(C) An equipment operator of a TRU must not cause or allow a TRU to operate within 1,000 feet of a residential area or school unless the cargo will be loaded or has been unloaded within 30 minutes.

(D) An owner of a vehicle, an off-road piece of equipment, or a TRU must ensure that:

- (1) The vehicle driver or equipment operator, upon employment and at least once per year

thereafter, is informed of the requirements of this Article, and of the consequences under this section, and the fleet owners terms of employment, of not complying with those requirements; and

- (2) Upon rental or lease of a vehicle or piece of equipment, notification is provided of the requirements of this subchapter;

(3) All complaints of non-compliance with, and enforcement actions related to the requirements of this subchapter are reviewed and remedial action is taken as necessary.

(E) A private property owner shall not allow a vehicle, an off-road piece of equipment or a TRU located on the owner's property to violate the provisions of this subchapter. A private property owner shall notify owners and operators of vehicles, off-road pieces of equipment, and TRUs entering the owner's private property of the requirements of this subchapter.

(Ord. 04-5, eff. 8-10-2004) Penalty, see § 71.99

§ 71.79 EXEMPTIONS.

(A) This subchapter does not apply to a vehicle or piece of equipment for the period or periods during which:

- (1) Idling is necessary while stopped:
 - (a) For an official traffic control device;
 - (b) For an official traffic control signal;
 - (c) For traffic conditions over which the driver has no control, including, but not limited to, stopped in a line of traffic, stopped at a railroad crossing, or stopped at a construction zone; or
 - (d) At the direction of a peace officer.
- (2) Idling is necessary to ascertain that the vehicle and/or the off-road equipment is in safe operating conditions and equipped as required by all provisions of law, and all equipment is in good working order, either as part of the daily vehicle inspection, or as otherwise needed;
- (3) Idling is necessary for testing, servicing, repairing or diagnostic purposes;
- (4) Idling is necessary for a period not to exceed three to five minutes (as per the recommendation of the manufacturer) to cool down a turbo charged heavy-duty vehicle before turning the engine off;
- (5) Idling is necessary to accomplish work for which the vehicle/equipment was designed, other than transporting goods, for example: operating a lift, crane, pump, drill, hoist, mixer or other auxiliary equipment other than a heater or air conditioner;
- (6) Idling is necessary to operate a life or other piece of equipment designed to ensure safe loading and unloading of goods and people;
- (7) Idling is necessary to operate defrosters, heaters, air conditioners, or other equipment to prevent a safety or health emergency, but not solely for the comfort of the driver or passengers;
 - (a) The only exception for driver comfort would be a vehicle driver that is required to have rest time by law. In this case, the driver may only idle at a designated rest area or truck stop and will not idle within 1,000 feet of a residential area or school;
 - (b) The only specific exception for passenger comfort would be vehicles with a passenger onboard with a disability or health condition that would be critically aggravated if the vehicle were not maintained at an adequate temperature.

(8) Idling is necessary solely to recharge a battery or other energy storage unit of a hybrid electric vehicle/equipment;

(9) Idling is necessary to operate equipment that runs intermittently;

(10) Alternative diesel fuel vehicles, or any Tier 2 4.8 g/bhp combined Nox and HMHC level;

(11) Idling is necessary in attainment portions of Placer County.

(B) Nothing in this subchapter allows idling in excess of other applicable laws, including but not limited to:

(1) Title 13 California Code of Regulations § 1226; requirement for leaving the driver's compartment when a pupil is aboard a school bus.

(2) Title 13 California Code of Regulations § 2480; requirements/restriction of idling of school buses.

(3) Cal. Vehicle Code § 22515; requirements for leaving a motor vehicle unattended.

(4) Any local ordinance or requirement as stringent as, or more stringent than, this chapter.

(Ord. 04-5, eff. 8-10-2004)

§ 71.80 ENFORCEMENT.

This subchapter may be enforced by the local air pollution control or air quality management district, and/or any peace officer as defined in Cal. Penal Code, Title 3, Chapter 4.5, §§ 830 *et seq.* and their respective agencies authorized representative(s).

(Ord. 04-5, eff. 8-10-2004)

§ 71.99 PENALTY.

(A) Any violation of this chapter for which a penalty is not provided shall be punished according to § 10.99 of this code.

(B) (1) For each violation of § 71.78, a driver of a vehicle, or an operator of an off-road piece of equipment or TRU will be first given a written warning. Subsequent violations will be subject to a civil penalty of \$50 and criminal penalties as provided by law.

(2) For each violation of § 71.78, an owner of a vehicle, off-road piece of equipment or TRU is subject to a written warning on the first offense, followed by a \$100 minimum civil penalty for a second offense, with a minimum civil penalty of \$200 for all future offenses and criminal penalties as provided by law. All penalties assessed under this section shall be deposited with the City of Auburn, regardless of whether another agency or entity first collects the penalties.

(Ord. 04-5, eff. 8-10-2004)

Cupertino Municipal Code

10.48.055 Motor Vehicle Idling.

Motor vehicles, including automobiles, trucks, motorcycles, motor scooters and trailers or other equipment towed by a motor vehicle, shall not be allowed to remain in one location with the engine or auxiliary motors running for more than three minutes in any hour, in an area other than on a public right-of-way, unless:

- A. The regular noise limits of Section 10.48.040 are met while the engine and/or auxiliary motors are running; or
- B. The vehicle is in use for provision of police, fire, medical, or other emergency services. (Ord. 1871, (part), 2001)

Fountain Valley Municipal Code

§ 6.28.147 Idling motor vehicles.

No person shall leave standing any motor vehicle, including refrigeration trailers, with engine idling or auxiliary motor running for in excess of ten minutes between the hours of ten p.m. and seven a.m. if the engine or motor noise disturbs the peace or quiet of any residential neighborhood or causes discomfort or annoyance to any reasonable person of normal sensitivity residing in the area. The driver, owner, registered owner and legal owner of the motor vehicle or refrigeration trailer shall each be guilty of the offense described herein. (Ord. 1156 § 1, 1990)

Palm Desert Municipal Code

10.98.010 Parking prohibitions and restrictions.

...

- C. While adjacent to a developed residential area within the city, the operator shall not idle the vehicles engine for longer than fifteen minutes. (Ord. 1025, 2002: Ord. 793 § 1 (part), 1996)

Placer County Code

Article 10.14 LIMITATION ON ENGINE IDLING

10.14.010 Findings and purpose.

The Placer County board of supervisors finds that:

A. Air pollution is a major public health concern in California. The Sacramento region is currently designated as non-attainment for the one-hour federal ozone standard, as well as the more stringent state ozone standard. Air pollution can cause or aggravate lung illnesses such as acute respiratory infections, asthma, chronic bronchitis, emphysema, and lung cancer. In addition to health impacts, air pollution imposes significant economic costs and negative impacts on our quality of life (nuisance).

B. Exhaust from vehicles (both on- and off-road) is a substantial source of ozone precursors in the Sacramento region. Vehicle exhaust is also a source of carbon monoxide, particulate matter, toxic air contaminants, and greenhouse gases. Although new engines have become cleaner due to improved emission control technologies; the slow turn over in their inventory and the number of miles/hours these vehicles idle each year is hindering progress in improving regional air quality.

C. Public agencies can play an important role in improving air quality by limiting the amount of time engines are allowed to idle within their jurisdiction. Public agencies have the responsibility to lead the effort to improve air quality by adopting ordinances that are cost effective in reducing ozone precursor emissions and toxic air contaminants. This article is based on and derived from the Sacramento Ozone Summit Model Engine Idling Ordinance.

D. A study of idling exhaust emissions conducted by the U.S. Environmental Protection Agency (EPA420-R-02-025, October 2002) indicates that a typical 1980s-2001 model year truck operating on diesel fuel emits one hundred forty-four (144) grams per hour of nitrogen oxide and eight thousand, two hundred twenty-four (8,224) grams per hour of carbon dioxide emissions and consumes about 0.82 gallons of diesel fuel while idling.

E. TIAx, a consultant for the Sacramento Metropolitan Air Quality Management District, estimated idling exhaust emissions from heavy heavy duty diesel trucks (HHDV), medium heavy duty diesel trucks (MHDV) and off road construction equipment to be 2.3 tons per day of nitrogen oxide emissions and .23 tons per day of reactive organic gas emissions. (Control Measures OFMS 52 and ONMS 45, April 2003). The maximum emissions reductions from full implementation of the Limitation on Engine Idling Ordinance in the Sacramento region was estimated to be 1.725 tons per day of nitrogen oxides emissions and .173 tons per day of reactive organic gas emissions (assuming a seventy-five percent (75%) compliance).

F. Under this article, a limitation on engine idling is established by the board of supervisors to discourage the idling of engines in the unincorporated Placer County. (Ord. 5271-B, 2003)

10.14.020 Definitions.

"Driver" means any person who drives, operates, or is in actual physical control of a vehicle.

"Emergency" means a sudden, urgent, usually unforeseen, occurrence.

"Equipment operator" means any person who is in actual physical control of a piece of off-road equipment.

"Gross vehicle weight rating" means the weight specified by the manufacturer as the loaded weight of a single vehicle.

"Commercial motor vehicle" means any on-road motor vehicle with a manufacturer's gross vehicle weight rating greater than twenty-six thousand (26,000) pounds or as defined in Motor Vehicle Code Section 15210(b).

"Idling" means the engine is running while the vehicle is stationary or the piece of off-road equipment is not performing work.

"Medium-duty vehicle" means any on-road motor vehicle with a manufacturer's gross vehicle weight rating of six thousand one to fourteen thousand (6,001 -- 14,000) pounds.

"Official traffic control device" means any sign, signal, marking or device, consistent with Section 21400 of the vehicle code, placed or erected by authority of a public body or official having jurisdiction, for the purpose of regulating, warning, or guiding traffic, but does not include islands, curbs, traffic barriers, speed humps, speed bumps, or other roadway design features.

"Official traffic control signal" means any device, whether manually, electrically, or mechanically operated, by which traffic is alternately directed to stop and proceed and which is erected by authority of a public body or official having jurisdiction.

"Off-road diesel equipment" means all non-road equipment with a horsepower rating of seventy (70) or greater.

"Transport refrigeration unit" or "TRU" means a refrigeration system powered by an engine designed to control the environment of temperature sensitive cargo. A TRU is a piece of off-road equipment regardless of its horsepower rating.

"Vehicle" means any on-road, self-propelled vehicle that is required to be registered and have a license plate by the Department of Motor Vehicles.

"Vehicle/equipment owner" means the registered owner, lessee, licensee, or bailee of any heavy- or medium-duty vehicle or piece of off-road equipment who operates or directs the operation of any such vehicle or equipment on either a for-hire or not-for-hire basis. (Ord. 5271-B, 2003)

10.14.030 Applicability.

There is established an article to be known as "Limitation on Engine Idling" that applies to the operation of all diesel fueled commercial vehicles over twenty-six thousand (26,000) lbs. gross vehicle weight rating, and all off-road diesel-powered equipment over seventy (70) horsepower rating, except as provided in Section 10.14.050. Additionally, this article applies to TRU engines as specified in subsection 10.14.040(C). (Ord. 5271-B, 2003)

10.14.040 Idling.

A. A driver of a vehicle:

1. Must turn off the engine upon stopping at a destination; and
2. Must not cause or allow an engine to idle at any location for more than five consecutive minutes.

B. An equipment operator of an off-road piece of equipment not identified in subsection A of this section must not cause or allow an off-road piece of equipment to idle at any location for more than five consecutive minutes.

C. An equipment operator of a TRU must not cause or allow a TRU to operate within one thousand (1,000) feet of a residential area or school unless the cargo will be loaded or has been unloaded within thirty (30) minutes.

D. An owner of a vehicle, an off-road piece of equipment, or a TRU must ensure that:

1. The vehicle driver or equipment operator, upon employment and at least once per year thereafter, is informed of the requirements in subsections 10.14.040(A)--(C), and of the consequences, under this section and the fleet owners terms of employment, of not complying with those requirements; and
2. Upon rental or lease of a vehicle or piece of equipment, notification is provided of the requirements in subsections 10.14.040(A)--(C); and
3. All complaints of non-compliance with, and enforcement actions related to, the requirements of subsections 10.14.040(A)--(C) are reviewed and remedial action is taken as necessary.

E. A private property owner shall not allow a vehicle, an off-road piece of equipment or a TRU located on the owner's property to violate subsections 10.14.040(A)--(C) respectively. A private property owner shall notify owners and operators of vehicles, off-road pieces of equipment, and TRUs entering the owner's private property of the requirements of subsections 10.14.040(A)--(C). (Ord. 5271-B, 2003)

10.14.050 Exemptions.

This article does not apply to a vehicle or piece of equipment for the period or periods during which:

A. Idling is necessary while stopped:

1. For an official traffic control device;
2. For an official traffic control signal;
3. For traffic conditions over which the driver has no control, including, but not limited to: stopped in a line of traffic, stopped at a railroad crossing, or stopped at a construction zone; or
4. At the direction of a peace officer;

B. Idling is necessary to ascertain that the vehicle and/or the off-road equipment is

in safe operating condition and equipped as required by all provisions of law, and all equipment is in good working order, either as part of the daily vehicle inspection, or as otherwise needed;

C. Idling is necessary for testing, servicing, repairing, or diagnostic purposes;

- D. Idling is necessary, for a period not to exceed three to five minutes (as per the recommendation of the manufacturer), to cool down a turbo-charged heavy-duty vehicle before turning the engine off;
- E. Idling is necessary to accomplish work for which the vehicle/equipment was designed, other than transporting goods, for example: operating a lift, crane, pump, drill, hoist, mixer, or other auxiliary equipment other than a heater or air conditioner;
- F. Idling is necessary to operate a lift or other piece of equipment designed to ensure safe loading and unloading of goods and people;
- G. Idling is necessary to operate defrosters, heaters, air conditioners, or other equipment to prevent a safety or health emergency, but not solely for the comfort of the driver or passengers;
 - 1. The only exception for driver comfort would be a vehicle driver that is required to have rest time by law. In this case, the driver may only idle at a designated rest area or truck stop and will not idle within one thousand (1,000) feet of a residential area or school.
 - 2. The only specific exception for passenger comfort would be a paratransit vehicle with a passenger on board with a disability or health condition that would be critically aggravated if the vehicle were not maintained at an adequate temperature.
- H. Idling is necessary solely to recharge a battery or other energy storage unit of a hybrid electric vehicle/equipment;
- I. Idling is necessary to operate equipment that runs intermittently;
- J. Alternative diesel fuel vehicles, or any Tier 2 4.8 g/bhp combined Nox and HMHC level;
- K. Idling is necessary in attainment portions of Placer County generally east of Donner Summit. (Ord. 5271-B, 2003)

10.14.060 Relationship to other laws.

Nothing in this article allows idling in excess of other applicable laws, including, but not limited to:

- A. Title 13 California Code of Regulations Section 1226;
- B. Title 13 California Code of Regulations Section 2480;
- C. Vehicle Code Section 22515; or
- D. Any local ordinance or requirement as stringent as, or more stringent than this article. (Ord. 5271-B, 2003)

10.14.070 Penalties.

- A. For each violation of subsections 10.14.040(A)--(C), a driver of a vehicle, or an operator of off-road piece of equipment or TRU is subject to a minimum civil penalty of fifty dollars (\$50.00) and to criminal penalties to the maximum extent provided by law.
- B. For each violation of subsection 10.14.040(D), an owner of a vehicle, off-road piece of equipment or TRU is subject to a warning on the first offense, followed by a one hundred dollar (\$100.00) minimum civil penalty for a second offense, with a

minimum civil penalty of two hundred dollars (\$200.00) for all future offenses and to criminal penalties to the maximum extent provided by law.

C. All fees collected through Section 10.14.070 or the penalty phase of this article shall be accrued in a vehicle replacement grant fund for annual application by commercial and off road vehicle operators. The air pollution control district will manage this fund. (Ord. 5271-B, 2003)

10.14.080 Enforcement.

This article may be enforced by the local air pollution control or air quality management district, and/or any peace officer as defined in California Penal Code, Title 3, Chapter 4.5, Sections 830 et seq. and their respective agencies' authorized representative. (Ord. 5271-B, 2003)

10.14.090 Effective date.

The operation and effective date of the ordinance codified in this article is January 1, 2004. (Ord. 5271-B, 2003)

Colorado

Municipal Codes

City of Aspen Municipal Code

13.08.110 Engine Idling.

(a) Except as hereinafter provided, it shall be unlawful for any person to idle or permit the idling of the motor of any stationary motor vehicle for a prolonged or unreasonable period of time determined herein to be five (5) minutes or more within any one (1) hour period of time.

(b) This section shall not apply when an engine must be operated in the idle mode for safety reasons including, but not limited to, the operation of cranes and fork lifts used in the construction industry.

(c) The time required by a diesel powered motor vehicle with a gross weight rating of ten thousand (10,000) pounds or more while operating in a stationary position to achieve a temperature of one hundred twenty (120) degrees Fahrenheit and an air pressure of one hundred (100) pounds per square inch, shall not be included in the computation of the five (5) minutes determined herein to be a prolonged or unreasonable period of time. The temperature and air pressure as indicated on the vehicle's gauges may be used for determining the diesel engine's temperature and air pressure.

(d) The time during which transportation vehicles are actively loading or discharging passengers shall not be included in the computation of the five (5) minutes determined herein to be a prolonged or unreasonable period of time. A transportation vehicle shall be defined for purposes of this section to mean motor vehicles designed to transport a minimum of sixteen (16) persons. (Ord. No 74-1992, § 1: Code 1971, § 11-2.70)

Denver Municipal Code

ARTICLE IV. MOBILE SOURCES

Sec. 4-43. Idling restriction.

(a) Effective July 1, 1990, no person shall allow a vehicle to idle for more than ten (10) minutes in any one-hour period unless:

(1) The ambient outside air temperature has been less than twenty (20) degrees Fahrenheit for the previous twenty-four-hour period; or

(2) The ambient outside air temperature is less than ten (10) degrees Fahrenheit.

(b) The idling restriction in subsection (a) shall not apply to emergency vehicles; to vehicles engaged in traffic operations; to vehicles which are being serviced; to vehicles that must idle to operate auxiliary equipment, including but not limited to pumps, compressors or refrigeration units; or to vehicles en route to a destination that are stopped by traffic congestion.

(Ord. No. 330-90, 6-4-90)

Greenwood Village Municipal Code

10.04.010 Model Traffic Code adopted.

A. The City hereby adopts by reference the 2003 edition of the Model Traffic Code for Colorado, promulgated and published by the Colorado Department of Transportation, Safety and Traffic Engineering Branch, 4201 East Arkansas Avenue, EP 700, Denver, CO 80222.

...

D. The City makes the following modifications to the 2003 edition of the Model Traffic Code:

1. The following new Part 3 is added to the Model Traffic Code:

*PART 3
EMISSIONS CONTROL*

...

304. Idling prohibited. It is unlawful for any person to operate or cause or knowingly permit to be operated in any residential district in the City, except on a state highway, any motor of a motor vehicle which weighs twelve thousand (12,000) pounds or more, manufacturer's gross vehicle weight, or any combination of motor vehicles towed by such motor vehicle, which remains stationary for a consecutive period longer than five (5) minutes.

Town of Johnstown Municipal Code

Sec. 8-44. Idling.

Motor vehicles that weigh more than ten thousand (10,000) pounds (mostly trucks) are hereby forbidden from idling longer than fifteen (15) minutes in any hour unless stopped due to traffic congestion. Moreover, said vehicles shall not idle within one hundred (100) feet of a residential area from 10:00 p.m. to 7:00 a.m. unless parked in the designated area.

Vail Town Code

5-1-7: NOISE PROHIBITED:

...

G.Motor Vehicle Noise:

...

3 Idling Engines:

a. It shall be unlawful for any person to idle or permit the idling of the engine of any bus, truck, or any motor vehicle of any kind whatsoever, for a period of time in excess of twenty (20) minutes within the Town limits.

b. Notwithstanding subsection G3a of this Section, it shall be unlawful for any person to permit any idling whatsoever of the engine of any unattended bus, truck or any motor vehicle, except for refrigeration vehicles, within the Lionshead Mixed Use 1, Lionshead Mixed Use 2, Commercial Core 1 or the Commercial Core 2 Zone Districts of the Town.

Winter Park Town Code

4-3-5: IDLING OF MOTOR VEHICLES:

A. The unreasonable and prolonged idling of motors of any motor vehicle of any kind whatsoever is hereby declared to be a nuisance and public safety and health hazard.

B. It shall be unlawful for any person to idle or permit the idling of the motor of any motor vehicle of any kind whatsoever for a prolonged and unreasonable period of time within the limits of the town at any time of the day or night.

C. Evidence that a motor vehicle has idled for a period of fifteen (15) minutes or longer shall be prima facie proof that said vehicle was idling for a prolonged and unreasonable period of time.

D. Any person convicted of a violation of this section shall be fined in an amount not to exceed three hundred dollars (\$300.00) or imprisoned for a term not to exceed ninety (90) days, or both, for each offense. (Ord. 334, Series of 2003)

Connecticut

State Codes

Connecticut General Statutes

§ 14-277. Operator's duties on stopping bus. Prohibition on idling of bus.

...

(b) The operator of any school bus shall not operate the engine of any school bus for more than three consecutive minutes when the school bus is not in motion except (1) when the school bus is forced to remain motionless because of traffic conditions or mechanical difficulties over which the operator has no control, (2) when it is necessary to operate heating, cooling or auxiliary equipment installed on the school bus when such equipment is necessary to accomplish the intended use of the school bus, including, but not limited to, the operation of safety equipment, (3) when the outdoor temperature is below twenty degrees Fahrenheit, (4) when it is necessary to maintain a safe temperature for students with special needs, (5) when the school bus is being repaired, or (6) when the operator is in the process of receiving or discharging passengers on a public highway or public road.

(c) Any person who violates any provision of this section shall, for a first offense, be deemed to have committed an infraction and for each subsequent offense shall be fined not less than one hundred dollars nor more than five hundred dollars.

Municipal Codes

Code of Town of Branford, CT

§ 189-6. Prohibited noise activities.

...

B. Truck idling. No person shall operate an engine or any standing motor vehicle with a weight in excess of 10,000 pounds manufacturer's gross vehicle weight (GVW) for a period in excess of 10 minutes when such vehicle is parked on a residential premises or on a Town road next to a residential premises.

Code of the Town of Mansfield

§ 134-7. Prohibited noise activities.

The following acts are deemed unlawful pursuant to the regulations contained herein. However, this enumeration shall not be deemed exclusive.

...

B. Truck idling. No person shall operate any standing motor vehicle with a weight in

excess of 10,000 pounds, manufacturer's gross vehicle weight (GVW), for a period in excess of 10 minutes when such vehicle is parked on or next to a residential premise.

Code of City of Norwalk

§ 44-10. Control of particulate emissions.

A. Visible emissions

- (1) No person shall cause or permit the emission of visible air pollutants with greater than twenty-percent opacity, except as permitted under the following sections.
- (2) A person may discharge air pollutants into the atmosphere from any source of emission for a period or periods aggregating not more than five minutes in any 60 minutes, provided that said air pollutants are of no greater than forty-percent opacity

C. Exceptions for uncombined water.

- (1) Where the presence of uncombined water, such as water vapor, is the only reason for the failure of an emission to meet the requirements of this regulation, then the provisions of this regulation shall not apply.
- (2) The following shall be exempt from the requirements of Subsection A(2):
 - (a) Antique automobiles over 30 years old.
 - (b) Mobile sources in the process of being repaired.
- (3) Emissions from stationary or idling mobile sources. No mobile-source engine shall be allowed to operate for more than three consecutive minutes when the mobile source is not in motion, except as follows:
 - (a) When a mobile source is forced to remain motionless because of traffic conditions or mechanical difficulties over which the operator has no control.
 - (b) When it is necessary to operate heating, cooling or auxiliary equipment installed on the mobile source when such equipment is necessary to accomplish the intended use of the mobile source.
 - (c) To bring the mobile source to the manufacturer's recommended operating temperature.
 - (d) When the outdoor temperature is below 20° F.
 - (e) When the mobile source is being repaired.
- (4) Subsections A(2) and C(3) shall not apply to aircraft, locomotives operating on

rails, vessels for transportation on water, lawnmowers, snowblowers and other small home appliances

...

§ 68-6. Prohibited activities.

A. General prohibition. It shall be unlawful for any person to make, continue or cause to be made or continued any loud, unnecessary and unreasonable noise.

B. The following activities are prohibited:

...

(2) Emissions from stationary or idling mobile sources. No mobile source engine shall be allowed to operate for more than three consecutive minutes when the mobile source is not in motion except as follows:

- (a) When a mobile source is forced to remain motionless because of traffic conditions or mechanical difficulties over which the operator has no control;
- (b) When it is necessary to operate heating, cooling or auxiliary equipment installed on the mobile source when such equipment is necessary to accomplish the intended use of the mobile source;
- (c) To bring the mobile source to the manufacturer's recommended operating temperature;
- (d) When the outdoor temperature is below 20° F;
- (e) When the mobile source is being repaired.

Windsor Code of Ordinances

Sec. 9-33. Prohibited noise activities.

The following activities are prohibited:

...

(2) *Truck Idling*: No person shall operate an engine of any standing motor vehicle with a weight in excess of ten thousand (10,000) pounds manufacturer's gross vehicle weight (GVW) for a period in excess of ten (10) minutes, when such vehicle is parked on a residential premise or on a town road next to a residential premise;

Delaware

Municipal Codes

Wilmington City Code

Sec. 37-6. Diesel-powered motor vehicles; idle standard.

(a) *Definitions.* In addition to the definitions set forth in section 37-1 of this chapter, for purposes of this section, the following words and terms shall have the following meanings, unless the context clearly indicates otherwise:

(1) *Diesel-powered motor vehicle* means a vehicle which is self-propelled by a compression ignition type of internal combustion engine and which is designed primarily for transporting persons or property on a public street or highway; for purposes of this section, passenger automobiles and motorcycles are excluded.

(2) *Idle* means the motor vehicle operating mode consisting of a nonloaded, throttled engine speed at the revolutions per minute specified by the manufacturer.

(b) *Standards.* No person shall cause, suffer, allow, or permit the engine of a diesel-powered motor vehicle to idle for more than three consecutive minutes if the vehicle is not in motion, except:

(1) A vehicle at the vehicle operator's place of business where the vehicle is permanently assigned may idle for 30 consecutive minutes; or

(2) A vehicle may idle for 15 consecutive minutes when the vehicle engine has been stopped for three or more hours.

(c) *Exceptions.* The provisions of subsection (b) of this section shall not apply to:

(1) Buses while discharging or picking up passengers;

(2) Vehicles stopped in a line of traffic;

(3) Vehicles whose primary or secondary power source is utilized in whole or in part for necessary and definitively prescribed mechanical operation other than propulsion, passenger compartment heating or air conditioning;

(4) Vehicles being or waiting to be examined by state or federal motor vehicle inspectors;

(5) Emergency vehicles in an emergency situation;

(6) Vehicles while being repaired;

(7) Vehicles while engaged in the process of connection, detachment or exchange of trailers; or

(8) Vehicles manufactured with a sleeper berth while being used, in a nonresidentially zoned area, by the vehicle's operator for sleeping or resting or in order to provide heat or air-conditioning.

(d) *Violations; penalties.* Violations of any provision of this section shall be punishable upon conviction in accordance with the provisions of section 37-186.

(Code 1968, § 37-20.1)

District of Columbia

District of Columbia Municipal Regulations

Title 20

Sec. 900.1

The engine of a gasoline or diesel powered motor vehicle, the engine of a public vehicle for hire, including buses with a seating capacity of twelve (12) or more persons, on public or private space shall not idle for more than three (3) minutes while the motor vehicle is parked, stopped, or standing, including for the purpose of operating air conditioning equipment in those vehicles, except as follows:

- (a) To operate private passenger vehicles;
- (b) To operate power takeoff equipment including, dumping, cement mixers, refrigeration systems, content delivery, winches, or shredders; or
- (c) To idle the engine for five (5) minutes to operate heating equipment when the ambient air temperature is thirty-two degrees Fahrenheit (32°F) or below.

Sec. 914.1

Each person who fails to comply with any of the provisions of this chapter, prevents any inspection authorized by this chapter, or keeps inaccurate records shall be punished by a fine not to exceed five thousand dollars(\$5,000).

Sec. 914.2

Each violation of, or failure to comply with, this chapter shall constitute a separate offense and the penalties described in §914.1 shall be applicable to each separate offense.

Georgia

Municipal Codes

Atlanta Code of Ordinances

Sec. 150-97. Restrictions for trucks, buses.

(a) *Time limit for trucks.* No person shall park or stand any truck or other freight-carrying vehicle, including any truck tractor, in excess of one-half ton capacity upon any public street or highway for longer than one hour at any time during the day or no person shall park any truck or other freight-carrying vehicle, including any truck tractor or their cabs, in excess of one-half ton capacity upon any public street or highway from 6:00 p.m. to 8:00 a.m. during standard time and from 8:00 p.m. to 7:00 a.m. during day light savings time.

(b) *Attendant required for certain trucks, buses.* No person shall stop or stand any truck or bus with a body more than eight feet wide or ten feet high on any street or public place without the driver or chauffeur being actually present and in charge thereof.

(c) *Time limit for idling.* No person shall stop or stand any truck or bus on any street or public place and idle for more than 15 minutes. A violation of this subsection shall, upon conviction, be punishable by a minimum fine of \$500.00. This limitation shall not apply under the following conditions:

(1) Emergency vehicles, utility company, construction and maintenance vehicles where the engines must run to perform needed work;

(2) Truck or bus is forced to remain motionless because of traffic conditions;

(3) Truck or bus is being used to supply heat or air conditioning necessary for passenger safety or comfort, and such truck or bus is being used for commercial passenger transportation or is a transit authority bus or school bus, in which idling shall be limited to a maximum of 25 minutes;

(4) If the ambient temperature is less than 32 degrees Fahrenheit, idling shall be limited to a maximum of 25 minutes; or

(5) Any vehicle, truck, bus, or transit authority bus in which the primary source of fuel is Natural Gas (CNG) or electricity shall be exempt from the idling limitations set forth in this section.

(Code 1977, § 13-2238; Ord. No. 2001-8, § 1, 2-13-01; Ord. No. 2002-84, § 1, 11-26-02)

Hawaii

State Codes

Hawaii Administrative Rules

§11-60.1-34

...

(c) No person shall cause, suffer, or allow any engine to be in operation while the motor vehicle is stationary at a loading zone, parking or servicing area, route terminal, or other off street areas, except:

- (1) During adjustment or repair of the engine at a garage or similar place of repair;
- (2) During operation of ready-mix trucks, cranes, hoists, and certain bulk carriers, or other auxiliary equipment built onto the vehicle or equipment that require power take-off from the engine, provided that there is no visible discharge of smoke and the equipment is being used and operated for the purposes as originally designed and intended. This exception shall not apply to operations of air conditioning equipment or systems;
- (3) During the loading or unloading of passengers, not to exceed three minutes; and
- (4) During the buildup of pressure at the startup and cooling down at the closing down of the engine for a period of not more than three minutes.

Illinois

State Codes

Illinois Compiled Statutes

§ 625 ILCS 5/11-1401. Unattended motor vehicles

Sec. 11-1401. Unattended motor vehicles. No person driving or in charge of a motor vehicle shall permit it to stand unattended without first stopping the engine, locking the ignition, removing the key from the ignition, effectively setting the brake thereon and, when standing upon any perceptible grade, turning the front wheels to the curb or side of the highway.

Municipal Codes

Cook County Environmental Control Ordinance

9.7 IDLING OF MOTOR VEHICLES

It shall be unlawful for any person to cause or permit the operation of the main engine of any motor vehicle when parked or standing, except for the following:

- (a) Whenever engaged in any rescue operations attendant to accident or other common disaster.
- (b) Whenever operation of the main power train is essential to a basic function as with, but not necessarily limited to, pre-mixed cement trucks, platform lift trucks, compactor refuse trucks, certain varieties of dump trucks and the like, while function is in action.
- (c) Whenever weather conditions justify the use of heating or air -conditioning systems for the welfare and safety of any occupants (or future passengers in the case of public vehicles stopped in turn around or other such waiting areas) or when such low temperatures prevail that the startup of public conveyances or service vehicles might not otherwise be feasible.
- (d) Whenever the need for operation of refrigeration equipment on trailers carrying perishable contents is necessary, but which then must conform with the appropriate boundary levels involved by location and most especially so when parked overnight in any district adjacent to occupied residences. In general when parked, the use of auxiliary power sources shall be subject to the same general caution regarding applicability of other noise level restrictions for operation of the main engine and when the vehicle is in motion shall be considered simply as a component of the overall resultant sound level as specified by Section 9.9(a) or in the case of private travel trailers with auxiliary air conditioners by 9.9(c) and these latter, even while legally parked are subject to the same lot line and zone noise level restrictions described above.
- (e) Whenever main or auxiliary engines are operated for emergency repairs, or when properly housed for professional maintenance (subject to appropriate boundary level restrictions) and the occasional maintenance such as cleaning and flushing of the radiator and associated circulation system and/or seasonal change of antifreeze, cleaning of the carburetor or the like of a personally owned auto by a private citizen.

Louisiana

Municipal Codes

City of New Orleans Code of Ordinances

ARTICLE II. BUSES

Sec. 122-52. Operating at idle.

The operator of a bus shall not allow buses to operate at idle on the streets of the city for a period longer than 20 minutes, except for situations beyond the operator's control and as otherwise provided for in chapter 162 and section 162-942.

(Code 1956, § 38-143)

Sec. 122-53. Operating at idle in the Garden District.

The operator of a bus shall not allow buses to operate at idle on the streets of the Garden District, which is bounded by St. Charles Avenue, Jackson Avenue, Louisiana Avenue and Magazine Street, for a period longer than ten minutes, except for situations beyond the operator's control.

(Code 1956, § 38-143.1)

...

ARTICLE IX. TOUR VEHICLES AND BUSES*

Sec. 162-941. Garden District tour bus route limitations.

...

(e) *Operation at idle.* It shall be unlawful for the operator of a bus to allow the bus to operate at idle for a period longer than ten minutes in the Garden District, except for situations beyond the operator's control.

(f) *Violations.* It shall be unlawful for the operator of any bus to violate the provisions of this section. Any person cited for violation of this section shall be subject to arrest or to the issuance of a summons or citation.

(Code 1956, § 12-249; M.C.S., Ord. No. 21480, § 5, 4-1-04)

...

(d) A properly licensed CPNC bus having an overall length in excess of 20 feet and not greater than 31 feet shall be permitted to use routes in the Vieux Carre as recommended by the director of the department of safety and permits and approved by the city council. The director of safety and permits shall, within 135 days of the adoption of this ordinance (Ordinance Number 17,558 M.C.S., as amended by Ordinance Number 17,727 M.C.S.), promulgate such rules and regulations, in compliance with section 2-1000 of the City Code, as enacted by Ordinance Number 17,611 M.C.S., necessary to enforce the provisions of this section. Such rules and regulations shall become effective, in accordance with section 4-107(3)(d) of the Home Rule Charter, only after approval by the

city council, after review and recommendation by the council's ground transportation committee or its successor. No bus operator while loading or unloading passengers shall be permitted to idle the bus engine longer than ten minutes. Further, no bus shall be allowed to stop in one place for a period in excess of 15 minutes.

...

Sec. 154-177. Schedule of fines and payments.

...

(b) The following is a minimum schedule of fines which may be paid in the violations bureau for a first offense, provided that the offense does not require a mandatory court appearance as set out in section 154-178 or in the schedule of fines; and that violations occurring simultaneously shall be considered as a single offense for the purpose of assessing fines.

Operation of bus at idle longer than 20 minutes	50.00
Operation of bus at idle longer than 10 minutes	30.00

Maine

Code of the Town of Bar Harbor

§ 194-38. Idling of motor vehicles. [Added 6-17-1997]

- A. Five-minute limitation. No person may cause or allow a motor vehicle to idle for more than five consecutive minutes while that vehicle is parked in any of the downtown areas during the time from May 1 to Columbus Day.
- B. Exceptions. The limitation set forth in the preceding subsection shall not apply to:
 - (1) Fire trucks, police cars, ambulances and other emergency vehicles while responding to an emergency call.
 - (2) Utility vehicles, including contractor's equipment, while engaged in the construction, maintenance or repair of utility facilities.
 - (3) Motor vehicles idling while in a traffic lane, as the result of congested traffic conditions beyond the driver's control (traffic jams).
 - (4) Refrigeration units of delivery vehicles.
- C. Prima facie evidence. The fact that a parked motor vehicle is idling in violation of this section shall be prima facie evidence that the unlawful idling was caused or allowed by the person in whose name that vehicle is registered.

Maryland

State Codes

Maryland Transportation Code

§ 22-402. Mufflers; prevention of noise; discharge of smoke; maximum period of idling

...

(c) Discharge of smoke; maximum period of idling. --

(1) No motor vehicle may be operated, nor may the owner or lessee of a motor vehicle permit it to be operated, on any highway in this State unless the engine power and exhaust mechanism is equipped, adjusted, and operated to prevent:

(i) The discharge of clearly visible smoke (comparable to smoke equal to or darker in shade than that designated as No. 1 of the Ringelmann Chart as published by the U.S. Bureau of Mines) in the exhaust emissions within the proximity of the exhaust outlet for more than 10 consecutive seconds; and

(ii) The discharge of smoke from any other part of the engine in such amounts and of such opacity as to partially obscure persons or objects from view.

(2) In this subsection, "smoke" means small gasborne and airborne particles, exclusive of water vapor, from a process of combustion in sufficient numbers to be observable.

(3) A motor vehicle engine may not be allowed to operate for more than 5 consecutive minutes when the vehicle is not in motion, except as follows:

(i) When a vehicle is forced to remain motionless because of traffic conditions or mechanical difficulties over which the operator has no control;

(ii) When it is necessary to operate heating and cooling or auxiliary equipment installed on the vehicle;

(iii) To bring the vehicle to the manufacturer's recommended operating temperature;
or

(iv) When it is necessary to accomplish the intended use of the vehicle.

(4) For a period of 1 year from July 1, 1971, this subsection shall be enforced by issuance of a warning. One year from July 1, 1971, it shall be enforced in the same manner as other violations of this section.

(5) This subsection does not apply to Class L (historic) vehicles.

Massachusetts

State Codes

Massachusetts General Law ch. 90

§ 16A. Unnecessary Operation of Engine of Stopped Motor Vehicle Prohibited; Exceptions; Penalty.

No person shall cause, suffer, allow or permit the unnecessary operation of the engine of a motor vehicle while said vehicle is stopped for a foreseeable period of time in excess of five minutes. This section shall not apply to (a) vehicles being serviced, provided that operation of the engine is essential to the proper repair thereof, or (b) vehicles engaged in the delivery or acceptance of goods, wares, or merchandise for which engine assisted power is necessary and substitute alternate means cannot be made available, or (c) vehicles engaged in an operation for which the engine power is necessary for an associate power need other than movement and substitute alternate power means cannot be made available provided that such operation does not cause or contribute to a condition of air pollution. Whoever violates any provision of this section shall be punished by a fine of not more than one hundred dollars for the first offense, nor more than five hundred dollars for each succeeding offense.

Municipal Codes

Cambridge Municipal Code

Section 10.17.100 Regulation of idling buses, trucks, and taxis and automobiles.

The Police Department shall promptly review and improve its enforcement of the statutory prohibitions against idling by busses, trucks and taxis and automobiles set forth at G.L., ch. 90, § 16A. Within two months of the effective date of the ordinance codified in this provision, the Commissioner of the Police Department shall report to the City Manager on the Department's implementation of this provision. (Ord. 1139 (part), 1992)

Code of the City of Chicopee

§ 260-30.1. Standing prohibited. [Added 6-18-1996 by Ord. No. 96-38]

- A. On the entire length of Thaddeus Street, trucks or any vehicles idling or found standing for more than five minutes will be subject to the following fines for each violation:
- (1) First offense: \$50.
 - (2) Second offense: \$100.
 - (3) Third offense: \$200.

City of Peabody Code of Ordinances

Sec. 19-90.3. Idling and overnight parking of trucks prohibited.

No person shall idle or park a truck upon any street or highway or part thereof as follows:

(1) Idling of trucks shall be adhered to in strict conformance to 310 CMR 7.11 regulated under the Department of Environmental Protection, Air Pollution Control, U Transportation Unit, (1) Motor Vehicles as follows:

a. All motor vehicles registered to operate in the Commonwealth of Massachusetts shall comply with pertinent regulations of the Registry of Motor Vehicles relative to exhaust and sound emissions.

b. No person shall cause, suffer, allow, or permit the unnecessary operation of the engine of a motor vehicle while said vehicle is stopped for a foreseeable period of time in excess of five (5) minutes.

(2) No person shall park overnight or idle its truck in excess of what is allowable under the parameters of 310 CMR 7.11 on any street or highway or part thereof in the City of Peabody within one hundred (100) yards of any residential property including property utilized for elderly housing as defined by Massachusetts State Statute.

(Ord. No. 11-99, § 2, 5-13-99)

Minnesota

Municipal Codes

Minneapolis Code of Ordinances

389.100. Prohibited acts. (a) The following acts are not allowed in the city and the causing thereof are prohibited:

...

(7) Idling of buses, trucks, tractors, truck-tractor, trailers and semitrailers, as those terms are defined in Minnesota Statute 168.011, while stopped, standing or parked in a residentially used area between the hours of 10:00 p.m. and 6:00 a.m. except as provided for under permit in section 389.70, in compliance with traffic signals or signs, at the directions of a police officer or while buses are in the act of loading or unloading passengers. This prohibition shall not apply to emergency vehicles of the police department, sheriff's office, fire department, nor to any public or private ambulances, nor to any public works or public utility vehicles where actually engaged in the performance of emergency or operational duties necessary to be performed by said public departments or public utilities, nor to any vehicle owned by or performing work for the United States of America or the State of Minnesota.

Owatonna City Code

Section 900:10. Idling of Engines in Residential Districts. No person shall have or allow a motor vehicle engine to idle in residential districts of this City, as defined in Ordinance No. 827 (Owatonna Zoning Ordinance), for longer than fifteen (15) minutes. No idling period shall be repeated at shorter intervals than five (5) hours.

St. Cloud Ordinance Code

Section 706:10. Idling of Engines. No person who has stopped or parked a vehicle at the edge or curb of that portion of West St. Germain Street from its intersection with 8th Avenue to its intersection with 10th Avenue shall idle or otherwise leave the engine of that vehicle running for a period of time in excess of 5 minutes.

Missouri

Municipal Codes

St. Louis City Revised Code

11.34.150 Restrictions of emission of visible air contaminants.

...

D. Motor vehicles, except for emergency vehicles, shall not operate in idle for more than ten (10) consecutive minutes.

Nevada

State Codes

Nevada Administrative Code

445B.576 Vehicles powered by gasoline or diesel fuel: Restrictions on visible emissions and on idling of diesel engines.

...

4. Except as otherwise provided in this subsection, a person shall not idle the engine of a diesel truck or a bus for more than 15 consecutive minutes. The provisions of this subsection do not apply to a diesel truck or a bus:

(a) For which the commission has issued a variance from the requirements of this subsection. A variance is not effective during an air pollution emergency as defined in the air quality plan for the State of Nevada.

(b) Which is an emergency vehicle.

(c) Used for the removal of snow.

(d) Used to repair or maintain other motor vehicles.

(e) Which is stopped because of traffic congestion while in transit on a highway, roadway or street.

(f) Which is idling while a repair or maintenance is being performed on it at a shop or facility for the repair and maintenance of motor vehicles.

(g) The emission from which is contained and treated by a method approved by the commission.

(h) The engine of which must idle to perform a specific task for which it is designed such as well drilling, trenching or hoisting. Such a diesel truck or a bus may not idle for more than 15 consecutive minutes during an air pollution emergency as defined in the air quality plan for the State of Nevada.

Municipal Codes

Clark County Air Quality Regulations

SECTION 45 - IDLING OF DIESEL POWERED MOTOR VEHICLES

45.1 Diesel Powered Motor Vehicle Idling

Except as otherwise provided in this subsection, a person shall not idle the engine of a diesel truck or a diesel bus for more than 15 consecutive minutes. The provisions of this subsection do not apply to a diesel truck or a bus:

- (a) For which the Clark County Air Pollution Control Hearing Board has issued a variance from the requirements of this subsection. A variance is not effective during an air pollution emergency episode stage declared by the Department of Air Quality and Environmental Management.
- (b) Which is an emergency vehicle.
- (c) Used to repair or maintain other MOTOR VEHICLES.
- (d) Which is stopped because of traffic congestion while in transit on a highway, roadway or street.
- (e) The EMISSION from which is contained and treated by a method approved by the CONTROL OFFICER.
- (f) The engine of which must idle to perform a specific task for which is it designed such as well drilling, trenching or hoisting. Such an engine may not idle for more than 15 consecutive minutes during an air quality emergency episode stage declared by the Department of Air Quality and Environmental Management.
- (g) Which is idling while maintenance procedures are being performed at a repair facility.

Washoe County District Board of Health Regulations

040.200 DIESEL ENGINE IDLING (Amended 12/15/93)

Except as otherwise provided in this subsection, a person shall not idle the engine of a diesel truck or a bus for more than 15 consecutive minutes. The provisions of this subsection do not apply to a diesel truck or a bus:

- A. Which is an emergency vehicle.
- B. Used for the removal of snow.
- C. Used to repair or maintain other motor vehicles.
- D. Which is traveling on a public right of way from one place to another.
- E. The engine of which must idle to perform a specific task for which it is designed such as well drilling, trenching or hoisting. Such a diesel truck or a bus may not idle for more than 15 consecutive minutes during an air pollution emergency episode stage declared by the Health Authority.
- F. When idling is necessary as part of a maintenance procedure performed at a repair facility.

New Hampshire

State Codes

New Hampshire Code of Administrative Rules

Env-A 1101.05 Operational Requirements for Diesel-Powered Motor Vehicles. The owner or operator of a diesel-powered motor vehicle shall comply with the following operational requirements unless specifically exempted from the operational requirements for diesel-powered motor vehicles:

- (a) When the temperature is above 0 °C (32 °F), a diesel engine shall not idle for more than 5 consecutive minutes;
- (b) When the temperature is -23 °C (-10 °F), 0 °C (32 °F), or anywhere in between the 2 temperatures, a diesel engine shall not idle for more than 15 consecutive minutes; or
- (c) When the temperature is below -23 °C (-10 °F), and where no nuisance is created, a diesel engine shall not be subject to idling restrictions.

Env-A 1101.06 Exemptions From the Operational Requirements for Diesel-Powered Motor Vehicles. The owner or operator of a diesel-powered motor vehicle shall be exempted from the operational requirements for diesel-powered motor vehicles when any one of the following conditions exists:

- (a) When a diesel-powered motor vehicle is forced to remain motionless because of traffic conditions over which the operator has no control;
- (b) When a diesel-powered motor vehicle is being used as an emergency motor vehicle;
- (c) When a diesel engine is providing power takeoff for refrigeration, lift gate pumps or other auxiliary uses, or supplying heat or air conditioning necessary for passenger comfort in those vehicles intended for commercial passenger transportation;
- (d) When a diesel-powered motor vehicle is being operated by a mechanic for maintenance or diagnostic purposes; or
- (e) When a diesel-powered motor vehicle is being operated solely to defrost a windshield.

New Jersey

State Codes

New Jersey Administrative Code

SUBCHAPTER 14. CONTROL AND PROHIBITION OF AIR POLLUTION FROM DIESEL-POWERED MOTOR VEHICLES

§ 7:27-14.3 General prohibitions

(a) No person shall cause, suffer, allow, or permit the engine of a diesel-powered motor vehicle to idle for more than three consecutive minutes if the vehicle is not in motion, except:

1. A motor vehicle at the vehicle operator's place of business where the motor vehicle is permanently assigned may idle for 30 consecutive minutes; or
2. A motor vehicle may idle for 15 consecutive minutes when the vehicle engine has been stopped for three or more hours.

(b) The provisions of (a) above shall not apply to:

1. A diesel bus while it is discharging or picking up passengers;
2. A motor vehicle stopped in a line of traffic;
3. A motor vehicle whose primary power source is utilized in whole or in part for necessary and definitively prescribed mechanical operation other than propulsion, passenger compartment heating or passenger compartment air conditioning;
4. A motor vehicle being or waiting to be examined by a State or Federal motor vehicle inspector;
5. An emergency motor vehicle in an emergency situation;
6. A motor vehicle while it is being repaired;
7. A motor vehicle while it is engaged in the process of connection or detachment of a trailer or of exchange of trailers; or
8. A motor vehicle, manufactured with a sleeper berth, while it is being used, in a non-residentially zoned area, by the vehicle's operator for sleeping or resting, unless the vehicle is equipped with a functional auxiliary power system designed in whole or in part to maintain cabin or sleeper berth comfort or to mitigate cold weather start-up difficulties.

(c) No person shall cause, suffer, allow or permit any emission control apparatus or element of design installed on any diesel-powered motor vehicle or diesel engine to be disconnected, detached, deactivated, or in any other way rendered inoperable or less effective, in respect to limiting or controlling emissions than it was designed to be by the original equipment or vehicle manufacturer, except for the purposes of diagnostics, maintenance, repair or replacement and only for the duration of such operations.

New Jersey Administrative Code

§ 7:27-15.8 Idle standard

(a) No person shall cause, suffer, allow, or permit the engine of a gasoline-fueled motor vehicle to idle for more than three consecutive minutes if the vehicle is not in motion.

(b) The provisions of (a) above shall not apply to:

1. Buses while discharging or picking up passengers;
2. Motor vehicles stopped in a line of traffic;
3. Motor vehicles whose primary and/or secondary power source is utilized in whole or in part for necessary and definitively prescribed mechanical operation other than propulsion, passenger compartment heating or air conditioning;
4. Motor vehicles being or waiting to be examined by State or Federal motor vehicle inspectors;
5. Emergency motor vehicles in an emergency situation;
6. Motor vehicles while being repaired;
7. Motor vehicles while engaged in the process of connection, detachment or exchange of trailers; or
8. Motor vehicles manufactured with a sleeper berth while being used, in a non-residentially zoned area, by the vehicle's operator for sleeping or resting.

Municipal Codes

Code of the City of Atlantic City

Article IV, Buses; Idling of Engines [Adopted 5-6-1982 by Ord. No. 38-1982]

§ 233-47.

Definitions.

For the purposes of this article, the following terms shall have the meanings indicated:

BUS -- Those vehicles capable of holding 15 or more passengers, which passengers originate outside of the City of Atlantic City.

STOPPING or STANDING -- Any cessation of movement of a bus, whether occupied or not, except in compliance with the directions of a police officer or traffic control sign or signal.

§ 233-48. Restrictions

Buses within the boundaries of the City of Atlantic City are not permitted to stop or stand with their engines running or idling in excess of five minutes, except when involved with loading or discharging passengers.

§ 233-49. Violations and penalties.

Any person violating any of the provisions of this article shall, upon conviction in the Municipal Court of the City of Atlantic City, be punished for each offense by a fine not to exceed \$500 or by imprisonment for any term not exceeding 90 days in the county jail, or in any place provided by the municipality for the detention of prisoners, or both, in the sole discretion of the Municipal Judge.

Township of Bernards Revised Ordinances

SECTION 3-13
Truck Idling

§ 3-13.1. Definitions.

The following words and terms, when used in this section, shall have the following meanings:

DIESEL-POWERED MOTOR VEHICLE shall mean a vehicle which is self-propelled by a compression-ignition-type of internal combustion engine and which is designed primarily for transporting persons or property on a public street or highway.

GASOLINE-FUELED MOTOR VEHICLE shall mean any motor vehicle equipped to be powered by a hydrocarbon fuel other than diesel fuel, but including alcohol fuels and hydrocarbon-alcohol fuel blends.

IDLE means the motor vehicle operating mode consisting of a nonloaded, throttled engine speed at the revolutions per minute specified by the manufacturer or at any other engine speed.

For purposes of this section, noncommercial passenger vehicles and motorcycles are excluded. (Ord. #1034)

§ 3-13.2. Idling Prohibited.

a. No person shall cause, suffer, allow or permit the engine of a diesel-powered or gasoline-fueled motor vehicle to idle for more than three consecutive minutes if the vehicle is not in motion, except:

1. A motor vehicle at the vehicle operator's place of business where the motor vehicle is permanently assigned may idle for 30 consecutive minutes; or
2. A motor vehicle may idle for 15 consecutive minutes when the vehicle engine has been stopped for three or more hours.

b. The provisions of Paragraph a above shall not apply to:

1. Buses while discharging or picking up passengers;
2. Motor vehicles stopped in a line of traffic;
3. Motor vehicles whose primary and/or secondary power source is utilized in whole or in part for necessary and definitively prescribed mechanical operation other than propulsion, passenger compartment heating or air conditioning;
4. Motor vehicles being or waiting to be examined by state or federal motor vehicle inspectors;
5. Emergency motor vehicles in an emergency situation;
6. Motor vehicles while being repaired;
7. Motor vehicles while engaged in the process of connection, detachment or exchange of trailers; or
8. Motor vehicles manufactured with a sleeper berth while being used, in a nonresidentially zoned area, by the vehicle's operator for sleeping or resting.

(Ord. #1034)

§ 3-13.3. Penalty

Any person violating any provision of this section shall, upon conviction thereof, be subject to a fine not less than \$100 nor greater than \$1,000 for each violation. (Ord. #1034)

Code of City of Cape May

Chapter 493: VEHICLES, IDLING OF

[HISTORY: Adopted by the City Council of the City of Cape May by Ord. No. 777 (Sec. 24-1 of the 1997 Revised General Ordinances). Amendments noted where applicable.]

GENERAL REFERENCES

Parking — See Ch. 362.

§ 493-1. Definitions.

As used in this chapter, the following terms shall have the meanings indicated:

IDLE — The motor vehicle operating mode consisting of a nonloaded, throttled engine speed at the revolutions per minute specified by the manufacturer.

MOTOR VEHICLE — All vehicles propelled otherwise than by muscular power, excepting motorized bicycles and such vehicles as run only upon rails or tracks.

PERSON — Corporations, companies, associations, societies, firms, partnerships and joint-stock companies as well as individuals, and shall also include all political subdivisions of this state or any agencies or instrumentalities thereof.

PUBLIC AND PRIVATE PROPERTY — All real estate within the City, including, inter alia, public and private parking lots, on which a motor vehicle may be physically located except for the public streets and highways within the City.

§ 493-2. Time limit for idling; exceptions.

- A. No person shall cause, suffer, allow or permit the engine of a gasoline-fueled or diesel-powered motor vehicle to idle for more than three consecutive minutes if the vehicle is not in motion.
- B. The aforesaid shall not apply to the following:
 - (1) A motor vehicle being operated upon the public highway which shall be governed by N.J.S.A. 39:3-70.2 and the Administrative Code Regulations adopted in connection therewith.
 - (2) Motor vehicles stopped due to a line of traffic.
 - (3) Emergency motor vehicles in an emergency situation.
 - (4) Motor vehicles being repaired.
 - (5) Motor vehicles in such circumstances as are deemed by the Chief of Police or his representative, designated in writing to require idling for a period in excess of three minutes due to the peculiar characteristics of the motor vehicle or the particular circumstance in which it is operating.

Code of the Borough of Closter

§ 183-13. Parking vehicles in prohibited areas; use of marked spaces; idling of commercial vehicles.

- A. No person shall park or leave standing a motor vehicle, whether attended or

unattended, on any of the roadways in a parking yard or parking place or in any place therein where parking is prohibited by notice given by a sign or otherwise.

- B. If parking spaces are provided, no person shall park or leave standing a motor vehicle, whether attended or unattended, except in a marked parking space. Such vehicle shall be parked properly within the lines of the parking space. [Added 6-12-1996 by Ord. No. 1996:715]
- C. No person shall leave or park a commercial motor vehicle over 10,000 pounds' gross weight on any street or in any parking yard within the Borough of Closter with the motor running or idling for more than 30 minutes. [Added 6-12-1996 by Ord. No. 1996:715]

Franklin Township Code

§ 253-190.11. Commercial vehicle parking.

A. Definition. As used in this chapter, a "commercial vehicle" shall mean an owner-operated commercial registered vehicle, or vehicle used for commercial purposes, with a gross vehicle weight (GVW) over 12,000 pounds. The standards of this section shall not apply to vehicles used in conjunction with an agricultural operation, recreational vehicles or to vehicles used for the transportation of children to school.

C. General standards.

...

(3) The idling of engines or operation of accessory equipment, such as refrigeration units, etc., shall be prohibited while a commercial vehicle is parked.

Code of the Borough of Hillsdale

Chapter 294: VEHICLES, IDLING OF

[HISTORY: Adopted by the Borough Council of the Borough of Hillsdale 12-8-1998 by Ord. No. 98-20. This ordinance provided that it shall take effect on 12-17-1998. Amendments noted where applicable.]

GENERAL REFERENCES

Air pollution — See Ch. 329.

§ 294-1. Purpose.

It is the policy of Hillsdale to prevent the air pollution caused by the idling of diesel powered motor vehicles that may jeopardize the health, welfare or safety of the citizens or degrade the quality of life.

§ 294-2. Definitions.

The following words and terms, when used in this chapter, shall have the following meanings, unless the context clearly indicated otherwise:

DIESEL BUS — Any diesel-powered autobus or motorbus of any size or configuration, whether registered in this state or elsewhere, that is designed or used for intrastate or interstate transportation of passengers for hire or otherwise on a public road, street or highway or any public or quasi-public property in this state, including but not limited to autobuses under the jurisdiction of the New Jersey Department of Transportation pursuant to Title 27 or 48 of the Revised Statutes; autobuses of the New Jersey Transit Corporation and its contract carriers that are under the inspection jurisdiction of the New Jersey Department of Transportation; autobuses under the authority of the Interstate Commerce Commission or its successor agency; school buses, as defined pursuant to N.J.S.A. 39:1-1; and hotel, casino, charter and special buses.

DIESEL ENGINE — A compression ignition type of internal combustion engine.

DIESEL-POWERED — Utilizing a diesel engine.

ELEMENT OF DESIGN — Any part of system on a motor vehicle or a motor vehicle engine pertaining to the vehicle's engine's certified configuration.

GROSS VEHICLE WEIGHT RATING or GVWR — The value specified by the vehicle manufacturer as the maximum loaded weight of a single or combination vehicle.

HEAVY-DUTY DIESEL VEHICLE — A diesel-powered motor vehicle, other than a diesel bus, that has a GVWR exceeding 8,500 pounds and is designed primarily for transporting persons or property.

IDLE — An operating mode where the vehicle engine is not engaged in gear and where the engine operates at a speed at the revolutions per minute specified by the engine or vehicle manufacturer.

MOTOR VEHICLE — All vehicles propelled otherwise than by muscular power, except motorized bicycles and such vehicles as run only upon rails or tracks.

PERSON — Any individual or entity and shall include, without limitation, corporations, companies, associations, societies, firms, partnerships and joint stock companies, and shall also include, without limitation, all political subdivisions of any states and any agencies or instrumentalities thereof.

QUASI-PUBLIC ROADWAY — Any roadway that, although under private ownership or control, is accessible to the public. This term shall include but not be limited to the New Jersey Turnpike, the Garden State Parkway and the Atlantic City expressway, but shall not include shopping mall roadways and parking lots, private business roadways, residential and nonresidential parking lots and private driveways.

§ 294-3. Prohibited acts; exceptions.

- A. No person shall cause, suffer, allow or permit the engine of a diesel-powered motor vehicle to idle for more than three consecutive minutes if the vehicle is not in motion, except that:
- (1) A motor vehicle at the vehicle operator's place of business where the motor vehicle is permanently assigned may idle for 30 consecutive minutes; or
 - (2) A motor vehicle may idle for 15 consecutive minutes when the vehicle engine has been stopped for three or more hours.
- B. The provisions of Subsection A above shall not apply to:
- (1) A diesel bus while it is discharging or picking up passengers.
 - (2) A motor vehicle stopped in a line of traffic.
 - (3) A motor vehicle whose primary power source is utilized in whole or part for necessary and definitely prescribed mechanical operation other than propulsion, passenger compartment heating or passenger compartment air conditioning.
 - (4) A motor vehicle being or waiting to be examined by a state or federal motor vehicle inspector.
 - (5) An emergency motor vehicle in an emergency situation.
 - (6) A motor vehicle while it is being repaired.
 - (7) A motor vehicle while it is engaged in the process of connection or detachment of a trailer or of exchange of trailers.
- C. No person shall cause, suffer, allow or permit any emission control apparatus or element of design installed on any diesel-powered vehicle or diesel engine to be disconnected, detached, deactivated or in any other way rendered inoperable or less effective, in respect to limiting or controlling emissions that it was designed to be by the original equipment or vehicle manufacturer, except for the purposes of diagnostics, maintenance, repair or replacements and only for the duration of such operations.

§ 294-4. Violations and penalties.

Any person who violates any provision of this chapter shall be subject to a penalty for each offense not more than \$2,500.

§ 294-5. Issuance of summons.

Local health officials, the Hillsdale Police Department and the local code enforcement office shall be empowered to enforce the provisions of this chapter.

§ 294-6. Other remedies.

No provision of this chapter shall be construed to impair any common law or statutory cause of action, or legal remedy therefrom, of any person for injury or damage arising from any violation of this chapter or from other law.

Princeton Township Code

Sec. 11-31.8. Prohibition on idling of diesel powered motor vehicles on township streets.

Pursuant to N.J.A.C. 7:27-14.3, no person may cause, suffer, allow or permit the engine of a diesel powered motor vehicle to idle for more than three consecutive minutes on streets within the Township of Princeton, if the vehicle is not in motion with the following exceptions:

- (a) The above provisions shall not apply:
 - (1) A diesel bus while it is discharging or picking up passengers;
 - (2) A motor vehicle stopped in a line of traffic;
 - (3) A motor vehicle whose primary power source is utilized in whole or in part for necessary and definitively prescribed mechanical operation other than propulsion, passenger compartment, heating or passenger compartment air-conditioning;
 - (4) A motor vehicle being, or waiting to be examined by a state or federal motor vehicle inspector;
 - (5) An emergency motor vehicle in an emergency situation;
 - (6) A motor vehicle is being repaired;
 - (7) A motor vehicle while it is being engaged in the process of connection or detachment of a trailer or of an exchange of trailer; or
 - (8) A motor vehicle manufactured with a sleeper berth while it is being used in a nonresidentially zoned area by the vehicle's operator for sleeping or resting, unless the vehicle is equipped with a functions auxiliary power system designed in whole or in part to maintain cabin or sleeper berth comfort or to mitigate cold weather startup difficulties;
 - (9) A motor vehicle at the vehicle operator's place of business where the motor vehicle is permanently assigned may idle for thirty consecutive minutes;
 - (10) A motor vehicle may idle for fifteen consecutive minutes when the vehicle engine has been stopped for three or more hours.

(b) Furthermore, no person shall cause, suffer, allow or permit any emission control apparatus or element of design installed on any diesel powered motor vehicle or diesel engine to be disconnected, detached, deactivated or in any other way rendered inoperable or less effective in respect to limiting or controlling emissions than it was designed to be by the original equipment or vehicle manufacturers, except for the purpose of diagnostics, maintenance, repair or replacement and only for the duration of such operation.

(Ord. No. 2004-9, § 1.)

Sec. 11-31.9 Idling of gasoline fueled motor vehicles on township streets.

Pursuant to N.J.A.C. 7:27-14.3, no person may cause, suffer allow or permit the engine of a gasoline fueled motor vehicle to idle for more than three consecutive minutes on streets within the Township of Princeton if the vehicle is not in motion with the following exceptions:

- (a) The above provisions shall not apply to:
 - (1) Buses while discharging or picking up passengers;
 - (2) Motor vehicles stopped in a line of traffic;
 - (3) Motor vehicles whose primary and/or secondary power source is utilized in whole or in part for necessary and definitively prescribed mechanical operation other than propulsion, passenger compartment heating or air conditions;
 - (4) Motor vehicles being or waiting to be examined by state or federal motor vehicle inspectors;
 - (5) Emergency motor vehicles in an emergency situation;
 - (6) Motor vehicles while being repaired;
 - (7) Motor vehicles while engaged in the process of connection, detachment or exchange of trailers; or
 - (8) Motor vehicles manufactured with a sleeper berth while being used in a nonresidentially zoned area by the vehicle's operator for sleeping or resting.

(Ord. No. 2004-9, § 2.)

Sec. 11-31.10. Penalty.

Any person who operated a motor vehicle or owns a motor vehicle which he/she permits to be operated on Township streets in violation of sections 11-31.8 or 11-31.9 of this Code shall be liable for a minimum penalty of a fine of not less than one hundred dollars and not more than one thousand dollars. In addition, any person violating sections 11-31.8 and 11-31.9 may be subject to imprisonment not exceeding ninety days or may be required to perform community service for a period not exceeding ninety days, all in accordance with section 1-6 of said Code. Any person who is convicted of violating sections 11-31.8 and 11-31.9 within one year of the date of a previous violation of the same provision and who was fined for that previous violation shall be sentenced by the court to an additional fine as a repeat offender. The additional fine imposed by the court upon a person for a repeated offense shall not be less than the minimum or exceed the maximum penalty set forth herein, but shall be calculated separately from the fine imposed for violation of sections 11-31.8 and 11-31.9.

(Ord. No. 2004-9, § 3.)

Code of the Borough of South River

§ 64-8. Idling of vehicles.

- A. No person shall cause, suffer, allow or permit the motor of a diesel-powered or gasoline-powered vehicle to be in operation for more than three consecutive minutes if the vehicle is not in motion, except where the ambient temperature is 32° F. or less, then the permitted period for idling shall be five consecutive minutes.
- B. The provisions of Subsection A shall not apply to:

- (1) Buses discharging or picking up passengers.
- (2) Vehicles stopped in a line of traffic.
- (3) Electric motor vehicles.
- (4) Emergency vehicles in performance of their prescribed function.
- (5) Vehicles whose primary and/or secondary power source is utilized in whole or in part for necessary and definitively prescribed mechanical operation other than propulsion.

New York

State Codes

New York Code of Rules and Regulations

SUBPART 217-3. IDLING PROHIBITION FOR HEAVY DUTY VEHICLES

§ 217-3.1 Applicability

This Part shall apply to all on-road heavy duty vehicles propelled by diesel fueled and nondiesel fueled engines excluding marine vessels. Heavy duty vehicle means a vehicle that has a GVWR exceeding 8,500 pounds and is designed primarily for transporting persons or properties.

§ 217-3.2 Prohibitions

No person who owns, operates or leases a heavy duty vehicle including a bus or truck, the motive power for which is provided by a diesel or nondiesel fueled engine or who owns, leases or occupies land and has the actual or apparent dominion or control over the operation of a heavy duty vehicle including a bus or truck present on such land, the motive power for which said heavy duty vehicle is provided by a diesel or non-diesel fueled engine, shall allow or permit the engine of such heavy duty vehicle to idle for more than five consecutive minutes when the heavy duty vehicle is not in motion, except as otherwise permitted by section 217-3.3 of this Subpart.

§ 217-3.3 Exceptions

The prohibitions of section 217-3.2 of this Subpart shall not apply when:

- (a) A diesel or nondiesel fueled heavy duty vehicle including a bus or truck is forced to remain motionless because of the traffic conditions over which the operator thereof has no control.
- (b) Regulations adopted by Federal, State or local agencies having jurisdiction require the maintenance of a specific temperature for passenger comfort. The idling time specified in section 217-3.2 of this Subpart may be increased, but only to the extent necessary to comply with such regulations.
- (c) A diesel or nondiesel fueled engine is being used to provide power for an auxiliary purpose, such as loading, discharging, mixing or processing cargo; controlling cargo temperature; construction; lumbering; oil or gas well servicing; farming; or when operation of the engine is required for the purpose of maintenance.
- (d) Fire, police and public utility trucks or other vehicles are performing emergency services.

(e) Trucks owned or operated by persons engaged in mining and quarrying are used within the confines of such person's property.

(f) A diesel fueled truck is to remain motionless for a period exceeding two hours, and during which period the ambient temperature is continuously below 25 degrees F.

(g) A heavy duty diesel vehicle, as defined in section 217-5.1(o) of this Part, that is queued for or is undergoing a State authorized periodic or roadside diesel emissions inspection pursuant to Subpart 217-5 of this Part.

(h) A hybrid electric vehicle, as defined in section 217-5.1(r) of this Part, idling for the purpose of providing energy for battery or other form of energy storage recharging.

(i) Heavy duty vehicles used for agricultural purposes on a farm.

(j) Electric powered vehicles.

Municipal Codes

Code of Town of Brighton

§ 104-2. Prohibited odors.

- A. No person shall cause or allow emissions of air contaminants or noxious odors to the outdoor atmosphere that are injurious to human, plant or animal life or to property or that unreasonably interfere with the comfortable enjoyment of life or property (for purposes of this chapter, a "prohibited odor").
- B. In addition to the general prohibition set forth in Subsection A above, the following shall be prohibited conduct in violation of this chapter, but the listing herein shall not be deemed to be exclusive.
 - (1) Any open burning in violation of § 73-1 of the Code of the Town of Brighton.
 - (2) The idling of any motor vehicle for a period of longer than 15 minutes within 10 feet of any lot line of a property.

Code of the Village of Bronxville

ARTICLE III, Idling of Motor Vehicles [Added 7-13-1992 by L.L. No. 2-1992]

§ 210-15. Operation of motor vehicle; idling of engine restricted.

A. No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking as defined in § 129 of the Vehicle and Traffic Law, standing as defined in § 145 of the Vehicle and Traffic Law, or stopping as defined in § 147 of the Vehicle and Traffic Law, unless the engine is used to operate a loading or unloading or processing device.

B. When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle while parking, standing or stopping.

§ 210-16. Enforcement.

Any parking enforcement officer or member of the Village of Bronxville Police Department is hereby authorized to issue and serve an appearance ticket as provided in Chapter 7, § 7-4, of this Code.

Code of Town of Clifton Park

§ 145-1. Findings and intent.

- A. Large vehicles, machines and equipment left idling or idling and unattended are a threat to the health, safety and welfare of the citizens of the Town of Clifton Park. Machines, equipment and vehicles of this nature include, but are not limited to, locomotives, tractor-trailer trucks and earth-moving equipment. When left idling for extended periods, these devices create a nuisance to Town residents in the form of excess noise and harmful exhaust fumes, and they also create an even greater danger to the public health and safety because of the possibility that they may be set in motion by passersby, including children.
- B. In order to protect and preserve the public health, safety and welfare, the Town of Clifton Park hereby restricts and proscribes the circumstances in which large machinery, equipment and vehicles may be left idling and unattended.

§ 145-2. Definitions.

As used in this chapter, the following terms shall have the meanings indicated:

EARTH MOVER — Mobile, mechanical equipment used in the excavation, displacement or transportation of earth.

IDLING— A circumstance in which an engine is running but not engaged in motion.

LOCOMOTIVE — A self-propelled engine, usually electric or diesel-powered, that pulls or pushes freight or passenger cars on railroad tracks.

MACHINERY AND EQUIPMENT — Locomotives, tractor-trailer trucks and earth movers.

TRACTOR-TRAILER TRUCKS — A truck having a cab and no body, used for pulling large vehicles such as vans or trailers.

UNATTENDED — A circumstance in which machinery or equipment is left idling with no operator present or in control of the equipment.

§ 145-3. Idling and unattended large machinery or equipment prohibited.

- A. It shall be unlawful for any person or entity to cause or to permit any locomotive, tractor-trailer truck or earth mover to idle for more than 10 minutes or to remain idling and unattended for more than five minutes.
- B. A law enforcement officer who observes idling machinery or equipment shall direct the operator to turn off the engine. In the event that the idling machinery or equipment is unattended, the officer shall turn off the engine. A private citizen who observes unattended machinery or equipment shall immediately report the circumstances to the police.

§ 145-4. Violations.

A law enforcement officer shall issue a citation for violation of this chapter in any instance in which such officer observes idling or unattended machinery or equipment. A citation may also be issued upon complaint of a citizen who observes idling or unattended machinery or equipment. A citation may be issued to an individual, a corporation or both. The prosecution of any citation shall be adjudicated before the Clifton Park Town Court. A violation of this chapter is classified as a misdemeanor.

§ 145-5. Penalties for offenses.

- A. Where an individual is adjudged guilty of a first violation of this chapter, the court may impose a fine not to exceed \$350 or imprisonment for a term of not less than 15 days nor more than one year, or both.
- B. Where an individual is adjudged guilty of a second violation of this chapter within a five-year period, the court may impose a fine not less than \$350 nor more than \$700 or imprisonment for a term of not less than 15 days nor more than one year, or both.
- C. Where an individual is adjudged guilty of a third violation of this chapter within a five-year period, the court may impose a fine not less than \$700 nor more than \$1,000 or imprisonment for a term of not less than 15 days nor more than one year, or both.
- D. Where a corporation is adjudged guilty of a violation of this chapter, the court may impose a fine of \$5,000.

Code of the Village of Flower Hill

§ 195-9. Idling of vehicles. [Amended 11-6-2000 by L.L. No. 3-2000]

Vehicles shall not be permitted to idle within the village in excess of two minutes.

City of Ithaca Code

§ 346-48. Vehicle idling

- A. Applicability. This section shall apply to all motor vehicles defined in Article 1

of the Vehicle and Traffic Law of the State of New York.

B. No person who owns, operates or leases a motor vehicle or who owns, leases or occupies land and has the actual or apparent dominion or control over the operation of a motor vehicle on such land shall allow or permit the engine of such motor vehicle to idle for more than five consecutive minutes when the motor vehicle is not in motion, except as otherwise permitted by Subsection C below.

C. Exceptions. The prohibitions of Subsection B of this section shall not apply when:

(1) The motor vehicle is forced to remain motionless because of traffic conditions over which the operator thereof has no control.

(2) Regulations adopted by federal, state or local agencies having jurisdiction require the maintenance of a specific temperature for passenger comfort. The idling time specified in Subsection B of this section may be increased, but only to the extent necessary to comply with such regulations.

(3) The engine is being used to provide power for an auxiliary purpose such as loading, discharging, mixing or processing cargo; controlling cargo temperature; construction; or farming, or operation of the engine is required for the purpose of maintenance.

(4) Fire, police and public utility trucks or other vehicles are actually performing emergency services.

D. Penalties for offenses. Any person who violates the provisions of this section shall be guilty of a violation and, upon conviction thereof, shall be punishable by a fine not to exceed \$250 or by imprisonment for not more than 15 days, or by both such fine and imprisonment.

Code of Village of Lawrence

§ 200-29.1. Bus idling. [Added 6-12-1991 by No. 2-1991]

No person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle for more than three minutes while parking, standing or stopping when the ambient temperature is in excess of 40° F. except while passengers are on board or while hoarding or discharging passengers.

Town of Mamaroneck Code

ARTICLE I Idling of Motor Vehicles [Adopted 5-4-2005 by L.L. No. 5-2005]

§ 219-1. Idling restricted.

A. No person shall allow, cause or permit the engine of any motor vehicle to run for more than five consecutive minutes while parking, standing, or stopping on public or private property in the unincorporated portion of the Town of Mamaroneck.

B. This section shall not apply to public utility companies, the United States of America, the State of New York, the County of Westchester, the Town of Mamaroneck, the Mamaroneck Union Free School District or the Villages of Larchmont, Mamaroneck

or Scarsdale. This section also shall not apply to any independent contractor engaged by any of the entities described in the preceding sentence while such independent contractor is carrying out the business activity for which it was engaged by one or more of the entities.

- C. This section shall not apply when the temperature in the Town of Mamaroneck is 40° F. or less.
- D. This section shall not apply in situations where a vehicle is exempt from the requirements of Subpart 217-3 of Title 6 of the New York Codes, Rules, and Regulations.

Village of Mamaroneck Code

§ 326-19. Engine idling prohibited.

- A. No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes on Village-owned property while parking, as defined in § 129 of the Vehicle and Traffic Law, while standing, as defined in § 145 of the Vehicle and Traffic Law, or while stopping, as defined in the Vehicle and Traffic Law, unless the engine is used to operate a loading, unloading or processing device.
- B. When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle while parking, standing or stopping on Village-owned property.

Code of Town of Milton

ARTICLE II Unattended or Idling Large Machinery and Equipment [Adopted 1-16-2002 by L.L. No. 1-2002]

§ 169-8. Findings and intent.

- A. Large vehicles, machines and equipment left idling or idling and unattended are a threat to the health, safety and welfare of the citizens of the Town of Milton. Machines, equipment and vehicles of this nature include, but are not limited to, locomotives, tractor-trailer trucks and earth-moving equipment. When left idling for extended periods, these devices create a nuisance to Town residents in the form of excess noise and harmful exhaust fumes, and they also create an even greater danger to the public health and safety because of the possibility that they may be set in motion by passersby, including children.
- B. In order to protect and preserve the public health, safety and welfare, the Town of Milton hereby restricts and proscribes the circumstances in which large machinery, equipment and vehicles may be left idling and unattended.

§ 169-9. Definitions.

As used in this article, the following terms shall have the meanings indicated:

EARTH MOVER — Mobile, mechanical equipment used in the excavation, displacement or transportation of earth.

IDLING — A circumstance in which an engine is running but not engaged in motion.

LOCOMOTIVE — A self-propelled engine, usually electric or diesel-powered, that pulls or pushes freight or passenger cars on railroad tracks.

MACHINERY AND EQUIPMENT — Locomotives, tractor-trailer trucks and earth movers.

TRACTOR-TRAILER TRUCKS — Trucks having a cab and no body, used for pulling large vehicles such as vans or trailers.

UNATTENDED — A circumstance in which machinery or equipment is left idling with no operator present or in control of the equipment.

§ 169-10. Prohibited acts; report of violations.

- A. It shall be unlawful for any person or entity to cause or to permit any locomotive, tractor-trailer truck or earth mover to idle for more than 10 minutes or to remain idling and unattended for more than five minutes.
- B. A law enforcement officer who observes idling machinery or equipment shall direct the operator to turn off the engine. In the event that the idling machinery or equipment is unattended, the officer shall turn off the engine. A private citizen who observes unattended machinery or equipment shall immediately report the circumstances to the police.

§ 169-11. Violations; issuance of citations.

A law enforcement officer shall issue a citation for violation of this article in any instance in which such officer observes idling or unattended machinery or equipment. A citation may also be issued upon complaint of a citizen who observes idling or unattended machinery or equipment. A citation may be issued to an individual, a corporation, or both. The prosecution of any citation shall be adjudicated before the Milton Town Court. A violation of this article is classified as a misdemeanor.

§ 169-12. Penalties for offenses.

- A. Where an individual is adjudged guilty of a first violation of this article, the court may impose a fine not to exceed \$350, or imprisonment for a term of not less than 15 days nor more than one year, or both.
- B. Where an individual is adjudged guilty of a second violation of this article within a

five-year period, the court may impose a fine of not less than \$350 nor more than \$700, or imprisonment for a term of not less than 15 days nor more than one year, or both.

- C. Where an individual is adjudged guilty of a third violation of this article within a five-year period, the court may impose a fine of not less than \$700 nor more than \$1,000, or imprisonment for a term of not less than 15 days nor more than one year, or both.
- D. Where a corporation is adjudged guilty of a violation of this article, the court may impose a fine of \$5,000.

Code of the City of New Rochelle

§ 312-33. Idling. [Added 9-21-2004 by L.L. No. 8-2004]

No person shall allow, cause or permit the engine of any motor vehicle to idle for more than five consecutive minutes while parking, standing, or stopping on public or private property in the City of New Rochelle, subject to the exceptions for heavy-duty vehicles set forth in Section 217.3 of Title 6 of the State of New York Codes, Rules, and Regulations.

New York City Administrative Code

34 RCNY § 4-08

§ 4-08 Parking, Stopping, Standing.

...

(p) Engine idling. (1) Idling of vehicle engines prohibited. Except as provided for buses in paragraph (p)(2) hereof, no person shall cause or permit the engine of any vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking, standing or stopping unless the engine is being used to operate a loading, unloading or processing device.

[SEE graphical material in printed version]

(2) Idling of bus engines prohibited. No person shall cause or permit the engine of any bus to idle at a layover or terminal location, whether or not enclosed, when the ambient temperature is in excess of forty (40) degrees Fahrenheit. When the ambient temperature is forty (40) degrees Fahrenheit or less, no person shall cause or permit any bus to idle for longer than three minutes at any layover or terminal location. For the purpose of this rule, at a layover or terminal location a bus engine shall not be deemed to be idling if the operator is running the engine in order to raise the air pressure so as to release the air brakes, provided however, that this shall not exceed a period of three minutes.

Code of Town of North Salem

ARTICLE VI, Idling of Vehicles [Added 8-25-1998 by L.L. No. 7-1998]

§ 220-31. Idling of motor vehicles restricted.

A. **Restrictions.** No person shall operate an engine of any standing motor vehicle for a period in excess of five minutes while parking as defined in § 129 of the Vehicle and Traffic Law, standing as defined in § 145 of the Vehicle and Traffic Law or stopping as defined in § 147 of the Vehicle and Traffic Law, unless the engine is used to operate a loading, unloading or processing device.

B. **Applicability.** The provisions of this chapter shall apply except when it is necessary to avoid conflict with other traffic or in compliance with the directions of a police officer or official traffic control device.

C. **Enforcement.** The provisions of this chapter shall be enforced by the Police Department of the Town of North Salem or other officials designated by the Town Board.

D. **Penalties for offenses.** Any person violating this chapter shall be guilty of a violation punishable by a fine of not more than \$500 or imprisonment not to exceed 15 days, or both such fine and imprisonment.

Code of Village of Northport

§ 289-1. Legislative findings.

The Board of Trustees hereby finds and determines that the excessive stationary idling of motor vehicles is detrimental to the health, safety and welfare of the village and its inhabitants in that it causes the release of unnecessary emissions of carbon monoxide and other pollutants into the atmosphere, the production of unnecessary noise and the waste of limited natural resources.

§ 289-2. Idling restrictions; exceptions.

- A. No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency vehicle, to idle for longer than five consecutive minutes when the ambient temperature is greater than 25° F. (-4° C.).
- B. Exceptions. The provisions of Subsection A shall not apply:
 - (1) When a vehicle is forced to remain motionless because of a traffic condition over which the operator has no control.
 - (2) When regulations adopted by federal, state or local agencies having jurisdiction require the maintenance of a specific temperature for passenger comfort, the idling limit specified in this section may be increased, but only to the extent necessary to comply with such regulations.
 - (3) When necessary to provide power for an auxiliary purpose, such as loading, discharging, mixing or processing cargo, controlling cargo temperature, construction, farming, or when the vehicle engine is being serviced.

- (4) When necessary for operation of mobile receiving and transmitter stations or mobile telephones.

§ 289-3. Penalties for offenses. Editor's Note: Amended at time of adoption of Code (see Ch. 1, General Provisions, Art. I).

Violations of any of the provisions of this chapter shall be punishable by a fine not to exceed \$250 and/or imprisonment for a term not to exceed 15 days, or both such fine and imprisonment, in the discretion of the court.

Code of Village of Nyack

§ 55-5.1. Idling prohibited. [Added 9-12-2002 by L.L. No. 4-2002]

- A. No commercial vehicle, bus, or other public omnibus shall be permitted to idle its engine for longer than five minutes on or along any street or public highway within the Village of Nyack.
- B. The penalty for a violation of this section shall be a fine not to exceed \$250.
- C. This section may be enforced by the Village of Nyack Parking Authority, the Clarkstown Police Department, or the Orangetown Police Department.

Code of Village of Port Chester

§ 319-26.4. Operation of motor vehicle; idling of engine restricted. [Added 11-1-1993 by L.L. No. 16-1993]

- A. No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking as defined in § 129 of the Vehicle and Traffic Law, standing as defined in § 145 of the Vehicle and Traffic Law or stopping as defined in § 147 of the Vehicle and Traffic Law, unless the engine is used to operate a loading, unloading or processing device.
- B. When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle while parking or standing.

Code of Town of Queensbury

ARTICLE I Idling in Residential Zones [Adopted 9-13-2004 by L.L. No. 7-2004]

§ 68-1. Title and authority.

The Queensbury Town Code is hereby amended by adding a new Article I of Chapter 68 entitled "Prohibition of Diesel Vehicles Idling in Residential Zones." It is adopted pursuant to Municipal Home Rule Law § 10.

§ 68-2. Legislative intent.

Concerns have been raised concerning idling of diesel vehicles, particularly in the Town's residential zones. Such activity can produce offensive odors and offensive noise which can harm the health, safety and welfare of residents. In addition, the activities prohibited in this article waste fuel which is a limited resource, cause environmental harm and are inappropriate in a residential zone. It is the finding of the Town Board that if a diesel vehicle needs to be warmed up or kept warm to run, then the licensed driver of the diesel vehicle shall use a block heater to keep the engine warm.

§ 68-3. Definitions.

For the purposes of this article, "diesel vehicle" shall mean any diesel-fueled vehicle requiring a CDL or higher license to drive. Notwithstanding the above, diesel vehicle shall not include emergency vehicles (e.g., fire or EMS), Town or other municipal or state vehicles operating in the course of their duties (including, without limitation, school buses) or utility company vehicles operating during an emergency, or delivery vehicles while making deliveries or making pickups at places, of business.

§ 68-4. Prohibition.

No person shall allow a diesel vehicle to run or idle its engine for any length of time while the diesel vehicle is unattended. "Unattended" means where the properly licensed driver of the diesel vehicle is not in the diesel vehicle's driver's seat and awake. This prohibition shall be effective in all residential zones of the Town. The driver and the registered owner of the diesel vehicle shall each be responsible for complying with this article.

§ 68-5. Enforcement.

Failure to comply with the provisions of this article shall be a violation and, upon conviction thereof, shall be punishable by a fine of not more than \$100 for the first offense. The second conviction hereunder shall be punishable by a fine of not more than \$200 or imprisonment for a period of not more than three days, or both. Any subsequent offense shall be punishable by a fine of not more than \$350 or imprisonment for a period of not more than seven days, or both. Each violation shall constitute a separate offense. The licensed driver of the diesel vehicle and, if different, the person in whose name the diesel vehicle is registered shall each be considered and responsible for each violation of this article and this article may be enforced against either one or both for each violation.

Code of the City of Rye

§ 45-1. Operation of motor vehicle; idling of engine restricted.

- A. No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking, as defined in § 129 of the Vehicle and Traffic Law, standing, as defined in § 145 of the Vehicle and Traffic Law, or stopping, as defined in § 147 of the Vehicle and Traffic Law, unless the engine is used to operate a loading, unloading or processing device.
- B. When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle while parking, standing or stopping.
- C. Violators of this chapter may be issued appearance tickets by police officers of the City of Rye. Such tickets shall be returnable in Rye City Court.

Code of the Village of Scarsdale

ARTICLE I, Motor Vehicle Engine Idling [Adopted 6-9-1992 by L.L. No. 2-1992]

§ 106-1. Idling of engines restricted.

No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking, as defined in § 129 of the Vehicle and Traffic Law, standing, as defined in § 145 of the Vehicle and Traffic Law, or stopping, as defined in § 147 of the Vehicle and Traffic Law, unless the engine is used to operate a loading, unloading or processing device.

§ 106-2. Bus engines restricted.

When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle while parking, standing or stopping.

§ 106-3. Violators issued appearance tickets.

Violators of this article may be issued appearance tickets by officers of the Scarsdale Police Department. Such appearance tickets shall be returnable in Scarsdale Village Court.

Code of Town of Somers

ARTICLE X Motor Vehicle Engine Idling [Adopted 7-9-1992 by L.L No. 10-1992]

§ 158-29. Restrictions.

- A. No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking as defined in § 129 of the Vehicle and Traffic Law, standing as defined in

§ 145 of the Vehicle and Traffic Law or stopping as defined in § 147 of the Vehicle and Traffic Law, unless the engine is used to operate a loading, unloading or processing device.

- B. When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle while parking, standing or stopping.

§ 158-30. Appearance tickets.

Violators of this article may be issued appearance tickets by officers of the Police Department of the Town of Somers. Such appearance tickets shall be returnable in local court.

§ 158-31. Penalties for offenses.

Any person violating the provisions of this article shall be guilty of an offense and shall be liable upon conviction for a fine not to exceed \$100 for each offense.

Laws of Suffolk County

§ 760-1017. Operation of motor vehicles.

A. No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency vehicle, to idle for longer than five consecutive minutes when the temperature is greater than 25° F. (-4° C.).

B. No diesel engine operated in the County of Suffolk shall emit a visible air contaminant of a shade of blue, black or gray of an opacity greater than 20% for a continuous period of more than five seconds.

C. No motor vehicle operated in the County of Suffolk shall emit a visible air contaminant of a shade of blue, black or gray of an opacity greater than 20% for a continuous period of more than five seconds.

D. Exceptions

(1) When a vehicle is forced to remain motionless because of a traffic condition over which the operator has no control.

(2) When regulations adopted by federal, state or local agencies having jurisdiction require the maintenance of a specific temperature for passenger comfort, the idling limit specified in this section may be increased, but only to the extent necessary to comply with such regulations.

(3) When necessary to provide power for an auxiliary purpose such as loading, discharging, mixing or processing cargo, controlling cargo temperature, construction, farming or when the vehicle engine is being serviced.

(4) When necessary for operation of mobile receiving and transmitter stations or mobile telephones.

Code of Ordinances of the Village of Tuckahoe

§ 21-86. Operation of motor vehicles idling of engines restricted.

- (a) No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking, as defined in § 129 of the Vehicle and Traffic Law, standing, as defined in § 145 of the Vehicle and Traffic Law, or stopping, as defined in § 147 of the Vehicle and Traffic Law, unless the engine is used to operate a loading, unloading or processing device.
- (b) When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus, as defined in § 104 of the Vehicle and Traffic Law, to idle while parking, standing or stopping.
- (c) A violation of this § 21-86 shall be punishable in accordance with § 1-7 of the Village Ordinances by a fine not to exceed \$250. (L.L. No. 7-1992, § 2)

...

§ 21-104.2. Prohibited uses

...

- (c) The provisions of § 21-86 of the Village Ordinances prohibiting the idling of engines of motor vehicles for more than three minutes shall be applicable to and enforced in parking lots.

Code of City of Yonkers

§ 109-88. Idling of engine. [Amended 2-11-1991 by G.O. No. 2-1991]

No person shall cause or permit the engine of a motor vehicle, other than a legally authorized emergency motor vehicle, to idle for longer than three minutes while parking, standing or stopping, unless the engine is used to operate a loading, unloading or processing device. When the ambient temperature is in excess of 40° F., no person shall cause or permit the engine of a bus as defined in § 104 of the Vehicle and Traffic Law to idle while parking, standing or stopping at any terminal point, whether or not enclosed, along an established route.

Ohio

Municipal Codes

Waynesville Code of Ordinances

§ 76.10 BUS STOPS AND TAXICAB STANDS; STANDING OR IDLING BUSES.

...

(B) Standing or idling busses.

(1) Buses shall be prohibited from standing and/or idling within the village, whether in the downtown section or otherwise, for a period in excess of 15 minutes.

(2) Violation of this division (B) may result in the issuance of a citation by the Police Department and summons into Mayor's Court.

(3) Each violation of this division (B) may result in a fine of up to \$50 per occurrence.

(4) For the purpose of this division (B), the definition of BUS shall be as defined in § 70.01.

(5) For the purpose of this division (B), the terms STANDING and IDLING shall be as defined under R.C. Title 45 or the administrative regulations adopted by the Ohio Department of Transportation, which definitions are expressly adopted herein. (Ord. 92-129, passed 10-19-92) Penalty, see 70.99

Oregon

Municipal Codes

City of Ashland Ordinance No. 2828

AN ORDINANCE ADDING SECTION 11.24.020.H TO THE
ASHLAND MUNICIPAL CODE TO PROHIBIT TRUCKS
AND BUSES FROM IDLING WHILE PARKED

THE PEOPLE OF THE CITY OF ASHLAND DO ORDAIN AS FOLLOWS:

SECTION 1. The following section is added to the Ashland Municipal Code as section 11.24.020.H:

SECTION 11.24.020 Prohibited parking. In addition to the provisions of the motor vehicle laws of Oregon prohibiting parking, no person shall park:

A. A vehicle upon a bridge, viaduct, or other elevated structure used as a street or within a street tunnel in this City, unless marked or indicated otherwise;

B. A vehicle in an alley except to load and unload persons or materials not to exceed twenty (20) consecutive minutes in any two (2) hour period;

C. A vehicle upon a street for the principal purpose of:

1. Displaying the vehicle for sale;
2. Washing, greasing, or repairing the vehicle except repairs necessitated by an emergency;
3. Selling merchandise from the vehicle except in an established marked place or when so authorized or licensed under the ordinance of this City;
4. Storage, or as junkage or dead storage for more than seventy-two (72) hours.

D. A vehicle upon any parkway except where specifically authorized;

E. A vehicle upon private property without the consent of the owner or person in charge of the private property;

F. A vehicle within any area marked off by yellow paint upon the street or upon the curb, except where specifically authorized by a traffic sign, (Ord. 1557 S13, 1968)

G. A vehicle or any part thereof upon a sidewalk or bicycle path. (Ord. 1971 S1, 1978)

H. Or stand or stop a truck or bus on a public street or in a public parking area with its engine running, if such engine emits exhaust fumes into the air. Vehicle engines shall be

turned off when loading and unloading passengers or merchandise. This subsection H shall not apply to:

1. An engine running for less than five minutes.
2. A vehicle in the moving traffic lane waiting to move with the normal flow of traffic,
3. An engine needed to operate equipment used to load or unload merchandise,
4. Trucks under 12,000 GVW and buses with a carrying capacity of fifteen passengers or less, or
5. Emergency vehicles, utility company, construction and maintenance vehicles, the engines of which must run to perform needed work.

The foregoing ordinance was first READ on the 16th day of April, 2002, and duly PASSED and ADOPTED this 7th day of May, 2002.

Barbara Christensen, City Recorder

SIGNED and APPROVED this 8th day of May, 2002

Alan DeBoer, Mayor

Reviewed as to form:

Paul Nolte, City Attorney

Pennsylvania

Municipal Codes

Allegheny County Health Department Rules and Regulations

§2105.91 School Bus Idling *{This Section added by September 8, 2004 Amendment, effective October 10, 2004.}*

a. Applicability. This Section applies to the operation of every heavy-duty diesel powered school bus.

b. General.

1. No school bus driver shall cause or allow the engine of any school bus subject to this section to idle prior to, during layover between, at the destination of, or at the conclusion of, any trip or route for more than five (5) consecutive minutes, except under the conditions described in Subsection c, below.

2. No school bus driver shall cause or allow the engine of a school bus subject to this section to be accelerated while idling, unless such action is taken in order to operate other equipment.

3. A school bus driver shall not park or idle a bus within 100 feet from a known and active school air intake system, unless the school district has determined that alternative locations block traffic, impair student safety or are not cost effective.

c. Exemptions. This section does not apply for the period or periods during which idling is necessary under the following circumstances:

1. Traffic Conditions.

A. For traffic conditions over which the driver has no control;

B. For an official traffic control device or signal; or

C. At the direction of a uniformed police officer or one of those persons authorized to direct traffic by the Vehicle Code, 67 Pa. Code §101.2.

2. Queuing at a School. Where the physical configuration of a school requires a queue of buses for the sequential discharge or pickup of students, and the queue of buses is actively engaged in the discharge or pickup of students.

3. Turbo-Charged Diesel Engine Cool Down or Warm Up. When the manufacturer's specifications require more time than the five minute limitation in §2105.91.b.1, above, to cool down or warm up a turbo-charged diesel engine.

4. Cold/Hot Weather.

A. If the outside temperature is less than 40°F, then idling is allowed for a period or periods aggregating not more than 20 minutes in any 60 minute period; or

B. If the outside temperature is greater than 75°F and the bus is equipped with air conditioning, then idling is allowed for a period or periods aggregating not more than 20 minutes in any 60 minute period.

5. Safety and Emergencies.

A. To ascertain that the school bus is in safe operating condition and equipped as required by all provisions of law, and all equipment is in good working order, either as part of the driver's daily vehicle inspection, or as otherwise needed;

B. To operate the flashing signal lamps and/or stop signal arm devices;

C. To operate defrosters, or other equipment to ensure the safe operation of the vehicle, or as otherwise required by federal or state motor carrier safety regulations, or other local requirements;

D. To operate a heater or an air conditioner of a bus that has, or will have, one or more children aboard with temperature sensitive disabilities;

E. To operate a lift or other piece of equipment designed to ensure safe loading, unloading,

or transport of persons with one or more disabilities; or

F. Use of school bus as an emergency vehicle.

6. Maintenance of Operations.

A. For testing, servicing, repairing, or diagnostic purposes; or

B. To recharge a battery or other energy storage unit of a hybrid electric bus.

d. Signage. Each school bus distribution center shall erect and maintain in a conspicuous location, a permanent sign(s) that is at least 12 inches by 18 inches in size indicating:

1. This school bus idling regulation in succinct language, and

2. The amount of money a violator will be fined.

e. Training. A motor carrier of a school bus shall ensure that the school bus driver, upon employment and at least once per year thereafter, is informed of the requirements of this Section and of the consequences of not complying with those requirements.

f. Penalties. Notwithstanding the provisions of Part I of this Article, violators of this Section are subject to:

1. A warning for the first offense;

2. A penalty of \$100 for the second offense; or

3. A penalty of \$500 for the third offense, and any subsequent offenses.

g. Enforcement. Notwithstanding any other provisions of this Article the prohibitions of this Section may be enforced by any municipal or local government unit having jurisdiction over the place where the idling occurs. Such enforcement shall be in accordance with the laws governing such municipal or local government unit and the Pa. Air Pollution Control Act. In addition, the Department may pursue the remedies provided by §2109.02 of this Article for any violation of this Section.

h. Relationship to Other Law. Nothing in this Section allows idling in excess of other applicable law, including, but not limited to any local ordinance or requirement as stringent as, or more stringent than, this Section.

Allegheny County Health Department Rules and Regulations

§2105.92 Diesel Powered Motor Vehicle Idling *{This Section added by June 13, 2005 Amendment, effective June 23, 2005.}*

a. Applicability. This Section applies to the operation of every heavy-duty diesel powered motor vehicle, except school buses.

b. General.

1. No driver shall cause or allow the engine of any heavy duty diesel powered motor vehicle subject to this section to idle prior to, during layover between, at the destination of, or at the conclusion of, any trip or route for more than five (5) consecutive minutes, except under the conditions described in Subsection c, below.

2. No driver shall cause or allow the engine of any heavy duty diesel powered motor vehicle subject to this section to be accelerated while idling, unless such action is taken in order to operate vehicle mounted accessory or service equipment.

c. Exemptions. This section does not apply for the period or periods during which idling is necessary for:

1. Traffic Conditions.

A. For traffic conditions over which the driver has no control;

B. For an official traffic control device or signal; or

C. At the direction of a uniformed police officer or one of those persons authorized to direct traffic by the Vehicle Code, 67 Pa. Code §101.2.

2. Boarding and Discharging Passengers.

A. When vehicles intended for commercial passenger transportation are boarding or discharging passengers; or

B. When vehicles intended for transporting people with disabilities are boarding or discharging passengers.

3. Queuing. When a vehicle, situated in a queue of other vehicles, must intermittently move forward to perform work or a service, and when shutting the vehicle engine off would impede the progress of the queue and be impracticable.

4. Turbo-Charged Diesel Engine Cool Down or Warm Up. When the manufacturer's specifications require more time than the five minute limitation in §2105.92.b.1, above, to cool down or warm up a turbo-charged diesel engine.

5. Cold/Hot Weather.

A. If the outside temperature is less than 40°F, then idling is allowed for a period or periods aggregating not more than 20 minutes in any 60 minute period; or

B. If the outside temperature is greater than 75°F and a vehicle is equipped with air conditioning, then idling is allowed for a period or periods aggregating not more than 20 minutes in any 60 minute period.

C. Notwithstanding subparagraphs A and B, in order to supply heat or air conditioning necessary for the comfort of passengers, a vehicle intended for commercial passenger transportation may idle for up to 10 minutes prior to passenger boarding and anytime passengers are onboard.

D. The Department may, upon request of an owner or manager of a bus terminal, approve alternate limits for warm-up of buses stored outdoors at the terminal when the temperature is below 40°F. Such plan shall include enforceable time limits that minimize bus idling.

6. Sleeping. When idling is necessary to power a heater, air conditioner, or any ancillary equipment during sleeping and resting in a truck cab or sleeper berth.

7. Safety and Emergencies.

A. To verify that the vehicle is in safe operating condition and equipped as required by all provisions of law, and all equipment is in good working order, either as part of the driver's daily vehicle inspection, or as otherwise needed;

B. To operate defrosters, or other equipment to ensure the safe operation of the vehicle, or as otherwise required by federal or state motor carrier safety regulations, or other local requirements; or

C. Use of vehicle as an emergency vehicle.

8. Operability and Maintenance.

- A. To provide power for vehicle mounted accessory or service equipment; or
- B. When being operated by a mechanic for testing, servicing, repairing, or diagnostic purposes.
- d. Penalties. Notwithstanding the provisions of Part I of this Article, violators of this Section are subject to:
 1. A warning for the first offense;
 2. A penalty of \$100 for the second offense;
 3. A penalty of \$500 for the third offense, and any subsequent offenses.
- e. Enforcement. Notwithstanding any other provisions of this Article the prohibitions of this Section may be enforced by any municipal or local government unit having jurisdiction over the place where the idling occurs. Such enforcement shall be in accordance with the laws governing such municipal or local government unit and the Pa. Air Pollution Control Act. In addition, the Department may pursue the remedies provided by §2109.02 of this Article for any violation of this Section.
- f. Relationship to Other Law. Nothing in this Section allows idling in excess of other applicable law, including, but not limited to any local ordinance or requirement as stringent as, or more stringent than, this Section.

Allegheny County Council Code of Ordinances

§ 56-1. Definitions.

As used in this article, the following terms shall have the meanings indicated:

DIESEL-POWERED MOTOR VEHICLE -- A self-propelled vehicle designed for transporting persons or property which is propelled by a compression-ignition type of internal-combustion engine.

HEAVY-DUTY -- Any motor vehicle with a gross vehicle weight of more than 8,500 pounds or with a passenger-carrying capacity of more than 12 persons.

IDLING -- The operation of an engine in the operating mode where the engine is not engaged in gear, where the engine operates at a speed at the revolutions per minute specified by the engine or vehicle manufacturer, or when the accelerator is fully released and there is no load on the engine.

MOTOR VEHICLE -- All vehicles propelled other than by muscular power except such vehicles as run only on rail or tracks.

PERSON -- Any individual, natural person, syndicate, association, partnership, firm, corporation, institution, agency, authority, department, bureau, or instrumentality of federal, state or local government or other entity recognized by law as a subject of rights and duties.

§ 56-2. Idling of diesel-powered motor vehicles.

The following provisions shall apply to the operation of heavy-duty diesel-powered motor vehicles:

- A. No person shall permit, cause, suffer or allow the engine of a heavy-duty diesel-powered motor vehicle to idle prior to, during layover between, or at the conclusion of, any trip or route for any period of time beyond that which is reasonably required to attain, or to secure from, normal operating conditions. The Board of Health shall promulgate rules and regulations, subject to the approval of the County Council, regarding a maximum allowable period of idling. Such rules and regulations shall also define exceptions to the maximum allowable period that consider extreme temperatures.
- B. No person shall permit, cause, suffer or allow the engine of a heavy-duty diesel-powered motor vehicle to be accelerated while idling unless such action is taken in order to operate other equipment.

...

§ 56-4. Exceptions to anti-idling policy.

The Board of Health shall promulgate rules and regulations, subject to the approval of the County Council, that exempt the owner or operator of a diesel-powered motor vehicle or diesel-powered locomotive when certain conditions exist. Those exemptions shall include, but not be limited to:

- A. When a diesel-powered motor vehicle or diesel-powered locomotive is forced to remain motionless because of traffic conditions over which the operator has no control;
- B. When a diesel-powered motor vehicle or diesel-powered locomotive is being used as an emergency vehicle;
- C. When a diesel engine is providing power takeoff for refrigeration, lift gate pumps or other auxiliary uses, or supplying heat or air conditioning necessary for passenger comfort in those vehicles intended for commercial passenger transportation;
- D. When a diesel-powered motor vehicle or diesel-powered locomotive is being operated by a mechanic for maintenance or diagnostic purposes; or
- E. When a diesel-powered motor vehicle or diesel-powered locomotive is being operated solely to defrost a windshield.

§ 56-5. Enforcement; violations and penalties.

It shall be the responsibility of the Department of Health to enforce this article and its provisions. Violators shall receive a warning for their first offense, a \$100 penalty for the second offense and a \$500 penalty for the third offense. The Department may make recommendations to Council regarding further enforcement mechanisms for this policy,

if appropriate.

Borough of Emsworth Code

§ 157-15. Idling of motor vehicles.

No person shall allow a vehicle to stand idle with the motor running for more than 10 minutes. Odors and fumes from motor vehicles shall not be permitted to affect the quiet enjoyment of adjoining residents. No vapors, malodorous gas or matter shall be permitted which is discernible on any adjoining property.

Township of Falls Code

§ 156-2. Prohibited odors.

- A. No person shall cause or allow emissions of air contaminants or noxious odors to the outdoor atmosphere that are injurious to human, plant or animal life or to property or that unreasonably interfere with the comfortable enjoyment of life or property (for purposes of this chapter, a "prohibited odor").
- B. In addition to the general prohibition set forth in Subsection A above, the following shall be prohibited conduct in violation of this chapter, but the listing herein shall not be deemed to be exclusive:
 - (1) Any open burning in violation of any Township ordinance.
 - (2) The idling of any motor vehicle for a period of longer than 15 minutes within 10 feet of any lot line of a property.

City of Philadelphia Air Management Regulations Ch. IX

**CONTROL OF EMISSIONS FROM MOBILE SOURCES
SECTION I. DEFINITIONS**

Air Contaminants - Any smoke, soot, flyash, dust, cinders, dirt, noxious or obnoxious acids, fumes, oxides, gases, mists, aerosols, vapors, odors, toxic or radioactive substances, waste, particulate, solid, liquid or gaseous matter, or any other materials in the outdoor atmosphere.

Diesel Powered Motor Vehicle - A self propelled vehicle designed for transporting persons or property which is propelled by a compression ignition type of internal combustion engine.

Gasoline Powered Motor Vehicle - A self propelled vehicle designed primarily for transporting persons or property which is propelled by a thermal ignition type of internal combustion engine, using gasoline as the fuel.

Motor Vehicle - All vehicles propelled other than by muscular power except such vehicles as run only on rails or tracks.

Opacity - The property of a substance which renders it partially or wholly obstructive to the transmission of visible light expressed as a percentage to which the light is obstructed.

Person - Any individual, natural person, syndicate, association, partnership, firm, corporation, institution, agency, authority, department, bureau, or instrumentality of Federal, State, or local government or other entity recognized by law as a subject of rights and duties.

...

SECTION III. IDLING OF DIESEL POWERED MOTOR VEHICLES

In addition to the emission limitations set forth in Section II, the following provisions shall apply to the operation of heavy-duty diesel powered motor vehicles:

A. No person shall permit, cause, suffer or allow the engine of a heavy-duty diesel powered motor vehicle to idle prior to, during layover between, or at the conclusion of, any trip or route for any period of time beyond that which is reasonably required to attain, or to secure from, normal operating conditions. The maximum allowable period of idling shall not exceed two (2) consecutive minutes or zero (0) for layovers, except under the following conditions:

1. The engine may be idled for a period of up to five (5) consecutive minutes when the ambient temperature is less than 32o F (0oC).
2. The engine may be idled for a period of up to twenty (20) consecutive minutes when the ambient temperature is less than 20oF (-7oC).
3. The engine may be idled for a period of up to twenty (20) consecutive minutes for buses equipped with air conditioning and non-openable windows and the ambient temperature is equal to or greater than 75o F.

B. No person shall permit, cause, suffer or allow the engine of a heavy-duty diesel powered motor vehicle to be accelerated while idling at any time.

For the purposes of this Section, the term "heavy-duty" shall apply to any motor vehicle with a gross vehicle weight of more than 8500 pounds or with a passenger carrying capacity of more than 12 persons.

SECTION IV. CIRCUMVENTION

No person shall build, erect, install or use any article, machine, equipment or other contrivance, the sole purpose of which is to dilute or conceal an emission without resulting in a reduction in total release of air contaminants to the atmosphere.

SECTION V. SEVERABILITY

The provisions of these Regulations are severable and if any provision, sentence, clause, section or part thereof shall be held illegal, invalid, unconstitutional or inapplicable to any person or circumstances, such illegality, invalidity, unconstitutionality or inapplicability shall not affect or impair any of the remaining provisions, sentences, clauses, sections or parts of the ordinance or their application to him or to other persons and circumstances. It is hereby declared to be the legislative intent that these regulations would have been adopted if such illegal, invalid, or unconstitutional provision, sentence, clause or part had not been included therein, and if the person or circumstance to which the ordinance or any part thereof is inapplicable had not specifically been exempted therefrom.

SECTION VI. EFFECTIVE DATE

Except as otherwise provided, this Regulation shall become effective upon adoption.

Code of Upper Providence Township

§ 169-23. Special purpose parking zones. [Amended 7-7-1997 by Ord. No. 363]

...

C. To promote the public health, safety and welfare by preventing unnecessary noise and air pollution, it is hereby declared that no person, corporation or entity shall leave any internal combustion engine-powered vehicle idling in any residential district as defined by the Zoning Ordinances of Upper Providence Township for more than a total of 60 minutes per day between the hours of 6:00 a.m. and 9:00 p.m. prevailing local time or for more than a total of 10 minutes between the hours of 9:00 p.m. and 6:00 a.m. For the purpose of this section, " idling " shall be defined as any running of an internal combustion powered vehicle while the vehicle is stationary, regardless of the revolutions per minute of the internal combustion engine at issue. Any person, corporation or entity violating this section shall be fined not less than \$75 and not more than \$600.

Rhode Island

Municipal Codes

Providence Code of Ordinances

Sec. 15-11. Residential parking/standing of certain commercial vehicles.

(a) For the purpose of this article, a commercial vehicle shall be defined as any vehicle in excess of 20 feet in length and any vehicle over 6,500 pounds gross vehicle weight.

(b) No commercial vehicle shall park or stand in any residentially zoned area unless engaged in the loading or off-loading of goods. At any such time the vehicle shall not be idling.

...

(e) Any person found in violation of any of these sections shall be subject to the penalties enumerated in section 1-10 herein, but in no case shall any person found in violation hereof be fined less than fifty dollars (\$50.00).

(f) (1) Any commercial vehicle otherwise in violation of this section and having an engine idling shall be deemed to be a hazard to the quality of life.

(2) Any commercial vehicle otherwise in violation of this section and in any way limiting, restricting or blocking the access of public safety vehicles in the course of their operation shall be deemed to be a hazard to public safety.

(3) Any commercial vehicle which, under this section, is deemed to be a hazard to the quality of life or a hazard to public safety shall be towed and the person violating subsection (f)(1) or (2) of this section shall be subject to a fine of not less than one hundred dollars (\$100.00).

(Ord. 2000, ch. 00-22, §§ 1--5, 8-11-00; Ord. 2002, ch. 02-11, §§ 1--6, 4-11-02)

South Carolina

Municipal Codes

City of Beaufort Code of Ordinances

Chapter 11. Tourism Management

Sec. 7-11027. Idling.

Idling of engines is allowed only while passengers are embarking onto or debarking from vehicles, not to exceed fifteen (15) minutes, with exceptions as noted below.

(Ord. No. O-13-04, 6-8-04)

Code of City of Charleston

ARTICLE VI. TOURING REGULATIONS GENERALLY

Sec. 29-239. Limitations on engine idling.

No buses may park with engines idling for more than five (5) minutes in residential areas.

(Ord. No. 1983-22, § 72, 5-10-83)

Texas

State Codes (only currently applicable to the following counties: Bastrop, Caldwell, Hays, Travis, Williamson and the following cities: Austin, Bastrop, Elgin, Lockhart, Round Rock, San Marcos)

Texas Administrative Code Title 30

RULE §114.510 Definitions

Unless specifically defined in the Texas Health and Safety Code, Chapter 382 (also known as the Texas Clean Air Act) or in the rules of the commission, the terms used in this subchapter have the meanings commonly ascribed to them in the field of air pollution control. In addition to the terms which are defined by Texas Health and Safety Code, Chapter 382; §3.2 of this title (relating to Definitions); §101.1 of this title (relating to Definitions); and §114.1 of this title (relating to Definitions), the following words and terms, when used in this subchapter, have the following meanings, unless the context clearly indicates otherwise.

- (1) Idle--The operation of an engine in the operating mode where the engine is not engaged in gear, where the engine operates at a speed at the revolutions per minute specified by the engine or vehicle manufacturer for when the accelerator is fully released, and there is no load on the engine.
- (2) Local government--A city, county, municipality, or political subdivision of the state.
- (3) Motor vehicle--Any self-propelled device powered by an internal combustion engine and designed to operate with four or more wheels in contact with the ground, in or by which a person or property is or may be transported, and is required to be registered under Texas Transportation Code, §502.002, excluding vehicles registered under §502.006(c).
- (4) Primary propulsion engine--A gasoline or diesel-fueled internal combustion engine attached to a motor vehicle that provides the power to propel the motor vehicle into motion and maintain motion.

RULE §114.511 Applicability

The provisions of §114.512 and §114.517 of this title (relating to Control Requirements for Motor Vehicle Idling; and Exemptions) are applicable only within the jurisdiction of a local government that has signed a Memorandum of Agreement with the commission to delegate enforcement of the provisions of this division to that local government.

RULE §114.512 Control Requirements for Motor Vehicle Idling

No person shall cause, suffer, allow, or permit the primary propulsion engine of a motor vehicle to idle for more than five consecutive minutes when the motor vehicle, as defined in §114.510 of this title (relating to Definitions), is not in motion during the period of

April 1 through October 31 of each calendar year.

RULE §114.517

Exemptions

The provisions of §114.512 of this title (relating to Control Requirements for Motor Vehicle Idling) do not apply to:

- (1) a motor vehicle that has a gross vehicle weight rating of 14,000 pounds or less;
- (2) a motor vehicle forced to remain motionless because of traffic conditions over which the operator has no control;
- (3) a motor vehicle being used by the United States military, national guard, or reserve forces, or as an emergency or law enforcement motor vehicle;
- (4) the primary propulsion engine of a motor vehicle providing a power source necessary for mechanical operation, not including propulsion, and/or passenger compartment heating, or air conditioning;
- (5) the primary propulsion engine of a motor vehicle being operated for maintenance or diagnostic purposes;
- (6) the primary propulsion engine of a motor vehicle being operated solely to defrost a windshield;
- (7) the primary propulsion engine of a motor vehicle that is being used to supply heat or air conditioning necessary for passenger comfort/safety in those vehicles intended for commercial passenger transportation or school buses in which case idling up to a maximum of 30 minutes is allowed;
- (8) the primary propulsion engine of a motor vehicle used for passenger transit operations in which case idling up to a maximum of 30 minutes is allowed;
- (9) the primary propulsion engine of a motor vehicle being used as airport ground support equipment; or
- (10) the owner of a motor vehicle rented or leased to a person who operates the vehicle and is not employed by the owner.

Utah

State Codes

Utah Code

§ 41-6a-1403. Motor vehicle left unattended -- Requirements

(1) A person operating or in charge of a motor vehicle may not permit the vehicle to stand unattended without:

- (a) stopping the engine;

Municipal Codes

Park City Municipal Corporation Municipal Code

9- 8- 3. DELIVERY VEHICLES IN THE MAIN STREET CORE.

All delivery vehicles parked on Main Street or Swede Alley shall observe the following restrictions:

...

(F) No delivery vehicle shall be parked with its engine left idling.

Salt Lake City-County Health Department Regulation #28

6.8 Vehicle Idling Limitation. No owner or operator of a diesel powered vehicle shall allow or permit such vehicle to remain in an idling mode or condition for a period of time exceeding fifteen (15) minutes.

6.8.1 Exemption to Idling Limitations. Vehicles may be exempted from the idling limitation requirements of Section 6.8 under the following conditions:

- (a). To supply power to a refrigeration unit for the purpose of cooling the contents of a trailer.
- (b). To provide heat or air conditioning to a sleeper unit of the vehicle.
- (c). Emergency vehicles.

6.8.2 Vehicles exempted from the Vehicle Idling Limitations as applicable under section 6.8.1 (a) and (b) of these regulations shall not remain in an idling condition for a period longer than fifteen (15) minutes if located within five hundred (500) feet of any residence.

Vermont

Municipal Codes

Burlington Code of Ordinances, Sec. 20-55

(e) No person shall leave idling for more than five (5) minutes any motor vehicle in any area of the city during the period from April 1 of every year to November 1 of the same year, except in the following instances:

(1) Motors used to run refrigeration units may be left idling to permit uninterrupted refrigeration;

(2) A motor vehicle may be left idling if necessary for the repair of that vehicle;

(3) This provision shall not apply to motor vehicles which must be kept idling in order to install, maintain or repair equipment or infrastructure.

(4) This provision shall not apply in any situation in which the health or safety of a driver or passenger requires the idling of the vehicle.

Virginia

State Codes

Virginia Administrative Code

ARTICLE 41. EMISSION STANDARDS FOR MOBILE SOURCES (RULE 4-41)

9 VAC 5-40-5670. Motor vehicles.

...

C. In commercial or residential urban areas, propulsion engines of motor vehicles licensed for commercial or public service use shall not be left running for more than three minutes when the vehicle is parked, unless the propulsion engine is providing auxiliary power for other than heating or air conditioning; except that:

1. Tour buses may idle for up to 10 minutes during hot weather in order to maintain power to the air conditioning system; and
2. Diesel powered vehicles may idle for up to 10 minutes to minimize restart problems.

Virginia Code

§ 46.2-1224.1. Local ordinances regulating certain parking; penalty

The governing body of any county having the county manager plan of government may by ordinance prohibit idling the engine of a bus for more than ten minutes when the bus is parked, left unattended, or is stopped for other than traffic or maintenance reasons.

Violators of such ordinance shall be subject to a civil penalty not to exceed fifty dollars, the proceeds from which shall be paid into the county's general fund.

The provisions of this section shall not apply to school buses or public transit buses.

Municipal Regulations

Arlington County Code

§ 14.2-2. Prohibition against parking of vehicles under certain conditions.

...

(f) No person shall idle the engine of a bus for more than ten (10) minutes when the bus is parked, left unattended, or is stopped for other than traffic or maintenance reasons. The provisions of this section shall not apply to school buses or public transit buses. Violators of this subsection shall be subject to a civil penalty of fifty dollars (\$50.00).

Fairfax County Code

Section 103-3-10. Mobile sources.

(a) *Prohibition of visible emissions from mobile sources.* No person shall cause or permit the emission of visible air contaminants from a mobile source of a density equal to or greater than twenty (20) percent opacity for longer than five (5) consecutive seconds after the operating engine of the mobile source has been brought up to operating temperature.

(b) *Prohibition of idling engines of mobile sources.* The operating engine of any mobile source shall not be left idling more than three (3) minutes after the mobile source has ceased to perform its designed work or function.

(27-78-103.)

Washington

Municipal Codes

Spokane Municipal Code

Title 15 Air Quality

Chapter 15.02 Carbon Monoxide Nonattainment Area Implementation Plan

Section 15.02.020 Idling Defined.

"Idling" means the running of an engine which supplies the motive power for a vehicle, when not for the purpose of moving the vehicle with the normal flow of traffic on a street or roadway. Idling does not include running the vehicle's engine while stopped at a traffic signal or waiting for the passage of other vehicles to permit safe entry into the flow of traffic.

Section 15.02.040 Excessive Idling Prohibited.

1. Excessive idling in the Central Business District Portion of the Nonattainment Area Prohibited.
Unless permitted by subsection (2) of this section it is unlawful for the operator of any vehicle powered by an internal combustion engine while located in the CBD portion of the nonattainment area to cause said engine to idle for a period longer than sixty seconds.
2. Idling Permitted.
To the extent necessary for the specified purpose, idling is permitted in the following cases:
 - a. police, sheriff or other law enforcement vehicles, including meter patrols, as required for safe and effective performance;
 - b. fire department vehicles, ambulances and other emergency vehicles when responding to an emergency or when the use of special equipment requires that the engines remain in operation;
 - c. construction or demolition equipment or other machinery when actually employed at the site of such work, and only to the extent necessary for efficient operation;
 - d. trucks, buses, or automobiles equipped with lift gates, winches, or other devices powered by take-offs from their engines, to the extent necessary to allow use of those devices provided; provided, however, that this subsection does not authorize idling solely for the purpose of operating any heating device, radio, power-assisted brakes, steering, seat adjustment or any luxury device not then required for safe operation of the vehicle;
 - e. vehicles participating in parades if licensed under Chapter 10.39;
 - f. buses or other mass transit vehicles while operated on a regular schedule;

- g. insofar as necessary for diagnosis, adjustment, or testing, vehicles undergoing repair or maintenance; but only when on the premises of the owner or person performing the work;
- h. any vehicle when required in an emergency for the protection of life, health, or property.

3. Notices.

The street director shall cause to be placed such notices as he may deem necessary to inform drivers of the prohibition against excessive vehicle idling; however, actual knowledge of the prohibition is not an element of an offense under this chapter.

Wisconsin

Municipal Codes

City of Madison Code

12.915 TRAFFIC ON STATE STREET RESTRICTED.

...

(3) Description of Area. This section applies to the 100 through 600 blocks of State Street running between West Mifflin Street and North Lake Street. When used in this section, the phrases “State Street” or “the area” shall include only the 100 through 600 blocks of State Street as described within this subsection. (Am. by Ord. 12,777, 3-13-01)

...

(7) Large Motor Truck Regulations.

...

(d) No operator of any large motor truck shall permit such vehicle to remain stationary with the main power train motor running for more than fifteen (15) minutes when the outside air temperature is between 40°F and 80°F. This subsection shall not apply to maintenance, construction or public utility vehicles. (Renum. by Ord. 12,777, 3-13-01)

Wyoming

Municipal Codes

Rawlins Municipal Code

Section 10.03.070 Vehicle Unattended

No person driving or in charge of a motor vehicle shall permit it to stand unattended without first stopping the engine, locking the ignition, removing the key from the ignition, effectively setting the brake thereon and, when standing upon any grade, turning the front wheels to the curb or side of the highway. Upon report of a vehicle running while unattended the police shall investigate and the person owning or in charge of the vehicle shall have five minutes from the time of the police arrival to shut off or move the vehicle. If after five minutes the vehicle has not been shut off or moved, a misdemeanor citation may be issued for violation of this section. No vehicle, even attended, shall remain idling while parked for more than 20 minutes. Any vehicle found in violation of this section may be towed at owner's expense. Upon repeat calls to the same property or for the same vehicle the police may immediately write a citation without waiting the five minutes.



Date: June 17, 2019

From: Erin Stokes, Cholera and Other Vibrio Illness Surveillance (COVIS) System Unit Coordinator *ES*
 Karen K. Wong, Lead, National Surveillance Team *KKW*
 Enteric Diseases Epidemiology Branch (EDEB)
 Division of Foodborne, Waterborne, and Environmental Diseases (DFWED)
 National Center for Emerging and Zoonotic Infectious Diseases (NCEZID)

Thru: Patricia M. Griffin, Chief *PMG*
 EDEB, DFWED, NCEZID

Subject: Recommendation for Interstate Shellfish Sanitation Conference (ISSC) *Vibrio vulnificus* Illness Review (VvIR) Committee procedure change

To: Ken Moore, ISSC Executive Director

Background

Currently, the ISSC VvIR committee uses information from COVIS to identify cases of *V. vulnificus* that meet criteria for a National Shellfish Sanitation Program (NSSP) Vv case. The current criteria include:

- Isolation of *V. vulnificus* from a clinical specimen, and
- Consumption of shellfish in the 7 days before illness began, specifically raw or undercooked, and
- The product was commercially harvested, and
- The case involved septicemia from consumption

As there is no “septicemia” variable on the COVIS case report form, determining septicemia requires committee members to interpret the available data for an individual case, making clinical judgments to infer whether an illness included septicemia. The clinical data collected on a COVIS case report form are brief. COVIS collects data for the purpose of public health surveillance, not clinical diagnosis. Furthermore, each state and interviewer collects and records data slightly differently. They may conduct patient or patient proxy interviews, review medical records, or interview a health care provider. Interviewers often do not have clinical training, and vary in their level of public health training.

Given the limitations of the surveillance data, it is important for scientific and procedural integrity that a case definition is unambiguous, with no need for clinical interpretation. The case definition should be clear enough that anyone can determine from the available data whether an illness fits the definition.

Septicemia is an outdated term that is no longer commonly used in medicine or public health. An alternative would be to classify cases by severity of disease. However, using only “severe” cases to reflect the magnitude of

risk from food is problematic, because the clinical severity of a case may depend on several factors other than the risk of the food itself, such as the patient’s age, underlying health conditions, access to healthcare, bacterial load ingested, and appropriateness of medical treatment. The clinical severity of individual cases does not necessarily reflect the risk to the public of *V. vulnificus*.

Recommendation

The most consistent approach to counting *V. vulnificus* cases for the purposes of monitoring risk would be to count all confirmed cases. Shifting to this broader case definition would require submission of a proposal to ISSC for consideration at the Biennial meeting.

In the interim, CDC recommends defining a case of severe *V. vulnificus* as illness in a person who had *V. vulnificus* infection confirmed by bacterial culture, and either of the following:

1. *V. vulnificus* was isolated from blood or a site that generally indicates spread of infection through the bloodstream, such as central nervous system sites, fluids from which bacteria cannot normally be cultured (excludes urine), and intra-abdominal sites (does not include stool). The table below lists sources that meet these criteria. This specimen source classification standard is used by the Enteric Diseases Epidemiology Branch across surveillance systems for multiple enteric conditions to define sources that likely reflect invasive infection.
2. Any of the following were indicated on the COVIS case report form:
 - a. Fever
 - b. Septic shock
 - c. Death
 - d. Any of the following sequelae: necrosis; or invasive procedure, such as surgery, amputation, skin graft, wound debridement, fasciotomy, or incision and drainage

CDC recommends immediate adoption of the suggested interim criteria. This will allow the VVIR committee to resume its case review work. Classification of cases according to these criteria should be reproducible regardless of who is doing the classification.

CDC plans to draft a proposal for conference consideration at the 2019 Biennial Meeting.

Table: Specimen sources that likely reflect invasive disease

Blood: Includes plasma and blood components
Vascular: Includes heart, heart valves, aorta, blood vessels
Lymphatic: Includes lymph, lymph nodes, thymus
Spleen: Includes spleen, splenic abscesses
Bone: Includes bone, bone marrow
Placenta and products of conception: Includes fetus, cord blood

<p>Nervous system</p> <ul style="list-style-type: none"> Cerebrospinal fluid (CSF) Other nervous tissue; includes brain abscess
Pleural fluid
Peritoneal fluid
Joint: includes synovial/joint fluid
Hepatobiliary: Gallbladder, bile, liver (includes abscesses)
Pancreas: Includes pancreas, pancreatic cysts, and abscesses
Reproductive: Ovary, fallopian tube, uterus (includes cysts and abscesses in these sites), pelvic abscesses, amniotic fluid
Kidney: Includes renal and perinephric abscess