PUBLIC HEALTH SERVICE U.S. FOOD AND DRUG ADMINISTRATION OFFICE OF FOOD SAFETY SHELLFISH AND AQUACULTURE POLICY BRANCH 5001 CAMPUS DRIVE

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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 SPE	CIALIST:		
OTHER OFFICIALS PRESENT:		TITLE:			
Items which do not conform are n	oted by:	Со	informity is noted by a " $$ "		
C – Critical K - Key O - Other N/A - Not Applicable					

	PART I – QUALITY ASSURANCE				
CODE	REF	ITEM			
CODE	KET	1.1 Quality Assurance (QA) Plan			
K	3, 6		1.1.1	Written Plan adequately covers all the following: (check $\sqrt{\text{those that apply}}$)	
K	3, 0		1.1.1	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 Ed	lucation	onal/Experience Requirements	
C	State's Human		1.2.1	In state/county laboratories, the supervisor meets the state/county	
	Resources			educational and experience requirements for managing a public health	
	Department			laboratory.	
K	State's		1.2.2	In state/county laboratories, the analyst(s) meets the state/county	
	Human Resources			educational and experience requirements for processing samples in a public	
	Department			health laboratory.	
С			1.2.3	In commercial/private laboratories, the supervisor must have at least a	
	USDA			bachelor's degree or equivalent in microbiology, biology, chemistry, or	
	Microbiology & EELAP			another appropriate discipline with at least two years of laboratory	
	C EEEM			experience.	
K	USDA		1.2.4	In commercial/private laboratories, the analyst must have at least a high	
11	Microbiology & EELAP		1.2	school diploma and shall have at least three months of experience in	
	& LLLIA			laboratory sciences.	
		1.3 W	ork A		
О	3, 6		1.3.1	Adequate for workload and storage.	
O	6		1.3.2	Clean and well lighted.	
0	6		1.3.3	Adequate temperature control.	
0	6		1.3.4	All work surfaces are nonporous and easily cleaned.	
	0			ory Equipment	
	4			• • •	
O K	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	
17			1 4 2	accuracy of 0.5 pH units.	
K	3		1.4.3	The pH meter is ca libra ted 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.	
		11			

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 on workdays.	
			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 on workdays. 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.	
О	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		
K	2	1.4.1/	Autopipettors are calibrated for the appropriate volumes used and checked	
	11	1 / 10	annually for accuracy. Results are recorded, and records are maintained.	
О	11	1.4.18	A centrifuge capable of holding 15 mL or 50 mL polypropylene tubes is	
		1.5 D	used.	
~			and Reference Solution Preparation and Storage	
<u>C</u>	11	1.5.1	All solvents and reagents used are ACS grade materials or better.	
О	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. Gela tin	
			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	1.5.1	3 Stock and working standard solutions are stored -10 °C or below.
С	5	1.5.1	
			date if not provided).
			ion and Transportation of Samples
О	4, 1	1.6.1	Shellstock are collected in clean, waterproof, puncture resistant containers.
K	4, 1	1.6.2	
С	4 1	1.6.3	shellstock, the harvest area, and time and date of collection. Immediately after collection, shellstock samples are placed in dry
	4, 1	1.0.3	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	
	ŕ		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
		1.5	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
			ation of Sample
C	4	2.1.1	1 1
			appropriate contingency plan for dealing with non-typical species of shellfish.
О	4	2.1.2	
О	4	2.1.3	1 , 5
О	4	2.1.4	The inside surfaces of the shells are rinsed with fresh water to remove sand and other foreign materials.
О	4	2.1.5	
			muscles and tissue connecting at the hinge.
С	4	2.1.6	
			opening.
О	4	2.1.7	1
77	4	210	layering for 5 minutes.
K 	4	2.1.8	
	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	
			(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 3000xg for 10 minutes.
С	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).
С	11	2.3.2	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	2.3.3	The working standard solution and crude sample extracts are brought to room temperature and thoroughly mixed before use. TMB is brought to room temperature in the dark prior to Assay Step 7.
С	11	2.3.4	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).
С	11	2.3.5	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 30 minutes, the liquid is poured from the plate, and all wells are rinsed 3 times with 300 µl PBS-Tween.
С	11	2.3.6	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.
С	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.
С	11	2.3.8	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.
С	11	2.3.9	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.
С	11	2.3.10	Assay Step 6: Reagent D is diluted by 800 (or as specified in kit instructions) in PGT, 100 μ l 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 μ l PBS-Tween, and one final time with 300 μ l PBS only to ensure no Tween remains on the plate.

C 11 2.3.11 Assay Step 7: Each well is filled with 100 µl of room TMB (3,3'5,5'-Tetramethylbenzidine) and incubated develops in the reference wells. The reaction is stopp µl of 0.5M sulfuric acid solution to each well, and the wells at 450 nm is measured in a microplate reaction.		
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the wells at 450 nm is measured in a microplate read		
K 11 2.3.12 Plates are covered with microplate sealing film during a	all incubation steps	
(Steps 1-6 above).		
C 11 2.3.13 Plates are protected from light by covering with alu	iminum foil during	
 color development (Step 7 above). The timing of the final step is reassessed with each new 	v lat afleit managemen	
	4 The timing of the final step is reassessed with each new lot of kit reagents and each new lot of TMB to ensure that reference well absorbance values	
	absorbance values	
fall into an appropriate range. 2.4 Quality Control		
	mosting the	
C 11 2.4.1 Acceptance of assay (plate) results is dependent on r following criteria:	meeting the	
a. Absorbance of standard reference wells (Amax)		
b. CV of raw absorbance of duplicate wells for stan		
the linear range of the assay (30-70% inhibition)) must be < 20%.	
C 11 2.4.2 Acceptance of individual sample results is dependen	nt on meeting the	
following criteria:		
a. CV of raw absorbance of duplicate wells for sam	ple dilutions used	
	20%.	
for quantitation (30-70% inhibition) must be < 2		
for quantitation (30-70% inhibition) must be < 20	nple dilutions within	
for quantitation (30-70% inhibition) must be < 200 b. CV of calculated concentrations of different same		
b. CV of calculated concentrations of different sam the linear range of the assay (30-70% inhibition)		
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- 10. Compendium of Methods for the Microbiological Examination of Foods, 3rd 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)

LABO	RATO	RY:	DATE OF EVALUATION:				
SHELLFISH LABORATORY EVALUATION CHECKLIST SUMMARY OF NONCONFORMITIES							
Page	Item	Observation	Documentation Required				
<u> </u>							
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LABORATORY STATUS					
LABORATORY	DATE				
LABORATORY REPRESENTATIVE:					
NEUROTOXIC SHELLFISH POISON (NSP or Brevetoxin) CO	OMPONENT: 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 brevetox	in (NSP) ELISA Component				
 Conforms Status: The NSP component of this Laboratory the following apply. a. No Critical nonconformities. b. and <6 Key nonconformities. c. and <12 Total nonconformities. Provisionally Conforms Status: The NSP component of the conforming to NSSP requirements if all of the following apara. the number of critical nonconformities is ≥ 1 but < 4. b. and < 6 Key nonconformities. c. and < 12 Total nonconformities. Does Not Conform Status: The NSP component of this laboratory 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. 	nis laboratory is determined to be provisionally ply.				
C. Laboratory Status (circle appropriate)					
•	Conforms				
Acknowledgement by Laboratory Director/Supervisor:					
All corrective Action will be implemented and verifying substantiati	ing documentation received by the Laboratory				
Evaluation Officer on or before					
Laboratory Signature:	Date:				
LEO Signature:	Date:				

NSSP Form 8-MARBIONC Brevetoxin ELISA Checklist, Rev. June 2024