**Laboratory Evaluation Checklist – Analysis for NSP (Mouse Bioassay)** 

# 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 301-436-2672

SHELLFISH LABORATORY EVALUATION CHECKLIST				
LABORATORY:				
ADDRESS:				
TELEPHONE:	FAX:	EMAI	IL:	
DATE OF EVALUATION:	DATE OF RE	PORT: LAST EVALUATION:		EVALUATION:
LABORATORY REPRESEN	TED BY:	TITLE:		
-				
LABORATORY EVALUATION OFFICER:		SHELLFISH SPECIALIST:		
		REGION:		
OTHER OFFICIALS PRESE	NT:	TITLE:		
Items which do not conform are noted by:				
C- Critical K - Key	O - Other	NA - Not Appli	cable	Conformity is noted by a " $$ "

Weighted	√	Item Description		
Code	V			
G		Quality Assurance (QA) Plan		
C		1. Written Plan adequately covers 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, calibration,		
-		maintenance, repair and performance. e. Laboratory safety.		
-		f. Internal performance assessment.		
G		g. External performance assessment.		
C		2. QA Plan is implemented.		
0		Work Area  Adagnets for workload and storage		
0		<ol> <li>Adequate for workload and storage.</li> <li>Clean and well lighted.</li> </ol>		
0		Clean and wen righted.     All work surfaces are nonporous and easily cleaned.		
C		4. A separate, quiet area with adequate temperature control is maintained for		
		acclimation and injection of mice.		
		Laboratory Equipment		
K		1. The differing sensitivities in weight measurements required by various steps in		
		the extraction procedure as well as the bioassay are met by the balances being		
		a. To determine sample weight, a sensitivity of at least 0.1 g at load of		
		100 g is required.		
		b. To determine the weight of the lipid extract and its subsequent volume		
		adjustment, a sensitivity of at least 10 mg at loads of 1 and 10 g is		
		required.		
		c. To determine the weight of the mice used in the bioassay, a		
0		sensitivity of 0.1 g at a load of 20 g is required.  2. The calibrations of the balances are checked monthly using NIST Class S or		
		ASTM Class 1 or 2 weights or equivalent. Records are maintained.		
K		3. The temperature maintained by the refrigerator is between 0 and 5°C.		
О		4. Refrigerator temperature is monitored at least once daily. Temperatures are		
		recorded and records are maintained.		
**		Reagents		
K		1. Concentrated (12N) HCl is used to acidify the homogenate.		
O C		2. Reagent grade NaCl is used in the extraction procedure.  2. Diathyl other purified for limit extraction is used for extracting limits from		
		3. Diethyl ether purified for lipid extraction is used for extracting lipids from the shellfish homogenates.		
C		4. Cottonseed oil (0.917 g/ml) or a solvent with a similar density (0.915 to 0.927		
		g/ml) is used as the toxin delivery system. Name of the solvent if substituted for		
		cottonseed oil. Specify density		
		Collection and Transportation of Samples		
0		1. Shellstock are collected in clean, waterproof, puncture resistant containers.		
K		2. Samples are appropriately labeled with the collector's name, the harvest area and		
K		the time and date of collection.  3. Immediately after collection, shellstock samples are placed in dry storage		
IX.		between 0 and 10°C until analyzed.		
K		4. Shellstock samples are analyzed within 24 hours of collection or		
		refrigerated unshucked until analyzed.		

T/		Defice and all stances of challete of does not arroad 40 hours		
K	5.	Refrigerated storage of shellstock does not exceed 48 hours.		
K	6.	If shellstock is refrigerated, only live animals are used in the analysis.		
K	7.	If shellfish are shucked in a location other than the laboratory, they must be		
	D	prepared according to steps 1-12 in "Preparation of Sample" section below.		
C		Preparation of Sample		
C	1.	At least 12 animals are used per sample.		
0	2.	The outside of the shell is thoroughly cleaned with fresh water.		
K	3.	Shellstock are opened by cutting the adductor muscles.		
C	4.	Shell liquor is discarded.		
О	5.	The inside of the shells is rinsed with fresh water to remove sand or other		
		foreign material.		
K	6.	Shellfish meats are removed from the shell by separating the adductor muscles		
		and tissue connecting at the hinge.		
K	7.	Damage to the body of the mollusk is minimized in the process of opening.		
K	8.	100 – 150 grams of meat are collected or all the available sample if there is less than		
		100 grams.		
О	9.	Shucked shellfish are drained on a #10 mesh sieve or equivalent without layering		
	- 10	for 5 minutes.		
K	10.	Pieces of shell and drainings are discarded.		
C	11.	Drained meats are blended at high speed until homogenous (60-120 seconds).		
C	12.	Shellfish homogenates are digested within 2 hours of blending.		
		ion of Sample		
K	1.	All glassware used is clean and properly washed with a succession of at least three		
		fresh water rinses, and a final distilled/deionized rinse to remove residual detergent.		
K	2.	100 grams (or entire sample amount if less than 100 grams is available) of		
		homogenized sample is weighted into a beaker.		
C	3.	1 ml of concentrated HCl and 5 g NaCl is added to the 100 gram homogenate		
		and thoroughly mixed. (For samples <100 g, add reagents to obtain final		
- C	4	concentrations of 0.12N HCl and 5% NaCl.)		
C	4.	The homogenate is brought to a boil and once $100 \pm 1^{\circ}$ C (sea level) is		
О	5.	reached, gently boil for 5 minutes.		
	3.	The beaker is covered with a watch glass or equivalent during boiling to prevent excessive evaporation.		
	6	<u> </u>		
0	6. 7.	The homogenate is boiled under adequate ventilation (fume hood).		
	/.	The boiled, acidified homogenate is cooled to room temperature or below in a		
	Extrac	refrigerator or in an ice bath.		
C				
C	1.	All steps in the extraction procedure which involve any manipulation of diethyl		
C	2.	ether are carried out under adequate.  100 ml of diethyl ether is added to the cooled, acidified homogenate in a		
	2.	stoppered centrifuge tube and shaken vigorously for 5 minutes.		
0	3.	Centrifuge tubes are vented frequently while being shaken and before being		
	3.	centrifuged to avoid accidents.		
C	4.	The content of the centrifuge tubes are centrifuged at 2000 rpm for 10 to		
	4.	15 minutes.		
C	5.	The clear upper ether phase is transferred to a large separatory funnel.		
C	6.	The contents of the centrifuge tube are extracted three additional times for		
	0.	a total of four times, each time with 100 ml of diethyl ether. The upper phases		
		are combined together in the separatory funnel (as in step 5).		
C	7.	The ether extract is transferred to a large, clean, dry pre-weighed beaker		
	/ '	(discard any emulsion or tissue that may have settled in the funnel.)		
C	8.	Ether is evaporated to dryness.		
C	9.	The final lipid residue is weighted and the weight is recorded.		
	7.	The linal upin residue is weighted and the weight is recorded.		

	Bioas	89V
С	1.	The volume of the lipid residue is adjusted by weight to 10 ml (9.17 g) per
	1.	100 g shellfish extracted using cottonseed oil. If a solvent with a density similar
		to cottonseed oil is used, the volume is adjusted to a weight 10 times the density
		of the solvent. Specify the weight to which the volume is adjusted to.
		of the softent. Speeny the weight to which the volume is adjusted to.
K	2.	A 25 gauge hypodermic needle is used for injection.
C	3.	Healthy male mice in the weight range of 17 to 23 grams from a stock colony
		are used for routine assays. Stock strain used Source of the
		mice
C	4.	Mice are allowed to acclimate for at least 24 hours prior to injection. In
		some cases up to 48 hours may be required. Typical length of the period of
		acclimation is
0	5.	Mice are weighed to the nearest 0.1 gram.
C	6.	The extract is completely mixed before it is injected.
C	7.	Mice are injected intraperitoneally with 1 ml of the lipid extract.
C	8.	A total of 5 mice are injected with undiluted or diluted extract as appropriate
		per sample in routine assays.
		a. The extract is not diluted when all test/assay mice survive beyond 110
		minutes of injection.
		b. The extract is diluted when 2 of 2 test mice or 3 of 5 assay mice survive
		for fewer than 110 minutes after injection
		c. When dilution is required, only dilutions which produce mean/median
		death times within 110 to 360 minutes of injection are used in the
C	9.	analysis. The time of completed injection is recorded.
C	10.	The time of completed injection is recorded.  Mice are continuously observed for at least 6 hours (360 minutes).
C	11.	If death occurs within the period of continuous observation, the time of
	11.	death to the nearest minute is noted by the last gasping breath.
K	12.	If mice survive the test, the time of death is recorded as ">" the period of
	12.	continuous observation.
	Calcu	llation of Toxicity
С	1.	The death time of each mouse is converted to mouse units (MU) using
		Table 8 in Recommended Procedures, 4th Edition.
0	2.	Table 8 is interpolated for death times between 110 and 360 minutes that are not
		listed in the Table.
K	3.	A weight correction in MU is made for each mouse injected using Table 8 in
		Recommended Procedures, 4 <sup>th</sup> Edition.
О	4.	Table 8 is interpolated to accommodate weights which are not listed.
С	5.	The death time for each mouse in MU is multiplied by a weight correction
		in MU to give the corrected mouse unit (CMU) for each mouse.
C	6.	The mean corrected mouse unit of the array of corrected mouse
		units (CMU) is used when all the mice injected with diluted or undiluted
		extract die during the period of continuous observation.
C	7.	The median corrected mouse unit of the array of corrected mouse units
		(CMU) is used when at least one mouse either survives the test or dies.
C	8.	The concentration of toxin is determined by the formula: Mean or median
		CMU x Dilution Factor x 10.
C	9.	When the time of death is known for certain for all mice injected, toxicity is
		determinate and the toxin concentration is reported as the number of mouse
		units per 100 grams of sample.

LABORATORY:			DATE OF EVALUATION:
SHELI	LFISH I	ABORATORY EVALUATION CHECKLIST	
SUMMARY OF NONCONFORMITIES			
Page	Item	Observation	Documentation Required
G			•

LABC	DRATORY STATUS					
LABC	DRATORY	DATE				
LABC	DRATORY REPRESENTATIVE:					
NEUR	ROTOXIC SHELLFISH POISON COMPONENT:					
Α.	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 NSP Component					
	<ol> <li>Does Not Conform Status The NSP component of this laboratory is not in conformity with NSSP requirements if:         <ul> <li>a. The total # of Critical nonconformities is ≥ 3 or</li> <li>b. The total # of Key nonconformities is ≥ 6 or</li> <li>c. The total # of Critical, Key and Other is ≥ 10</li> </ul> </li> <li>Provisionally Conforms Status: The NSP component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but &lt; 3</li> </ol>					
C.	Laboratory Status (circle appropriate)					
	Does Not Conform Provisionally Conforms	Conforms				