.12 Evaluation of Laboratories by State Shellfish Laboratory Evaluation Officers Including Laboratory Evaluation Checklists

SHELLFI	PUBLIC HEALT FOOD AND DRUG A OFFICE OF FOO SH AND AQUACUL 5100 PAINT BRANG COLLEGE PARK, I L. 240- 402-2151/2055	ADMINISTRAT OD SAFETY TURE POLICY CH PARKWAY MD 20740-3835	BRANCH		
SHELLFISH	H LABORATORY E	VALUATION C	HECKLIST		
LABORATORY:					
ADDRESS:					
TELEPHONE:	FAX:				
EMAIL:					
DATE OF EVALUATION:	DATE OF R	EPORT:	LAST EVALUATION:		
LABORATORY REPRESENTED BY	· ·	TITLE:	I		
	•				
LABORATORY EVALUATION OFF	ICER:	SHELLFISH S	PECIALIST:		
OTHER OFFICIAL C PRECENT		REGION:			
OTHER OFFICIALS PRESENT:		TITLE:			
Items which do not conform are noted C- Critical K - Key O - Other NA- Not A		onformity it note	ed by a "√"		
Check the applicable analytical method	ds:				
Multiple Tube Fermentation T	echnique for Seawater	(APHA)[PART I	II]		
Multiple Tube Fermentation T	echnique for Seawater	using MA-1 [PA	RT II]		

		Membrane l	Filtratio	on Techni	que for Seawater using mTEC [PART II]			
		Multiple Tu	be Fer	mentation	Technique for Shellfish Meats (APHA)[PART III]			
		Standard Pla	ate Cou	unt for She	ellfish Meats [PART III]			
		Elevated Te	mperat	ture Colife	orm Plate Method for Shellfish Meats [PART III]			
		Male Specific Coliphage for Soft-shelled Clams and American Oysters [PART III]						
PAF	RT 1	l - QUALI	TY A	SSURA	NCE			
CO	DE	REF.			ITEM			
			1.1 Q	uality As	ssurance (QA) Plan			
				1.1.1	Written Plan (Check those items which apply.)			
					a. Organization of the laboratory.			
					b. Staff training requirements.			
k	7	8, 11			c. Standard operating procedures.			
	•	0, 11			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.			
(7	8		1.1.2	QA Plan Implemented.			
k	ζ.	11	П	1.1.3	The Laboratory participates in a proficiency testing program annually.			
					Specify Program(s)			
			1.2 E	ducation	al/Experience Requirements			
C	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	ζ.	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	7	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	ζ	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 V	Work Ar				
(8,11		1.3.1	Adequate for workload and storage.			
k		11		1.3.2	Clean, well-lighted.			
k	<u> </u>	11		1.3.3	Adequate temperature control.			
()	11		1.3.4	All work surfaces are nonporous, easily cleaned and disinfected.			
k	ζ	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 La	aborator	y Equipment			
()	9		1.4.1	To determine the pH of prepared media, the pH meter has a standard accuracy of 0.1 units.			
C)	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.
О	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.
С	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.
С	11	1.4.14	Temperature of the waterbath is maintained at 44.5 ± 0.2 °C under all loading conditions.
С	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.
С	4	1.4.19	All working thermometers are appropriately immersed.
С	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 Devises (RTDs) and Platinum Resistance Devices (PTDs).
С	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
С	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 $\leq \pm 0.05$ °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.
О	11		1.4.25	Appropriate pipet aids are available and used to inoculate samples. Mouth pipetting is not permitted.
		1.5 Lab	oware a	nd Glassware Washing
О	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.
О	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.
			1.5.6	Pipettes used to inoculate the sample deliver accurate aliquots, have unbroken
С	9			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.
С	2		1.5.9	An alkaline or acidic detergent is used for washing glassware/labware.
С	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 Ste	rilizati	on and Decontamination
K	9		1.6.1	Autoclave(s) are of sufficient size to accommodate the workload.
О	8		1.6.2	Routine autoclave maintenance is performed and the records are maintained.
С	11, 30		1.6.3	The autoclave provides a sterilizing temperature of $121\pm2^{\circ}C$ 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
			1.6.6	Working autoclave thermometers are checked against the autoclave standards
K	1			thermometer at 121°C yearly.
	-			Data of last sheet
			1.6.7	Date of last check Method Spore strips/suspensions appropriate for use in an autoclave media cycle are
IZ.	1.1		1.0.7	used monthly according to manufacturer's instructions to evaluate the
K	11	Ш		effectiveness of the sterilization process. Results are recorded and the records
0	1.1		1 6 0	maintained.
0	11		1.6.8	Heat sensitive tape is used with each autoclave batch.
			1.0.9	Autoclave sterilization records including length of sterilization, total heat exposure time and chamber temperature are maintained.
K	11, 13			
		_		Type of record: Autoclave log, computer printout or chart recorder tracings.
			1 6 10	(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.
17	0		1.6.11	A thermometer capable of determining temperatures accurately in the range of
K	9			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
		<u> </u>	1.6.13	the hot-air sterilizing oven during use. Spore strips/suspensions are used quarterly to evaluate the effectiveness of the
K	11	ΙШ	1.0.13	sterilization process in the hot-air oven. Records are maintained.
K	11	П	1.6.14	Reusable sample containers are sterilized for 60 minutes at 170°C in a hot-air
11	11		1 6 15	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
С	1			each lot received. Results are recorded and the records maintained.
K	9	П	1.6.17	Reusable pipettes are stored and sterilized in aluminum or stainless steel
			1.6.18	canisters. Reusable pipettes (in canisters) are sterilized in a hot-air oven at 170°C for 2
K	9	Ш	1.0.10	hours.
С	2		1.6.19	The sterility of reusable pipettes is determined with each load sterilized. Results
<u> </u>			1.600	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.
			1.6.21	Hardwood applicator transfer sticks are properly sterilized.
K	18			
			1.600	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.
0	12		1.6.23	Spent broth cultures and agar plates are decontaminated by autoclaving for at
0	13			least 30 minutes before conventional disposal.
		1.7 Me		paration
K	3, 5		1.7.1	Media is commercially dehydrated except in the case of medium A-1 which
	[1	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.
О	11		1.7.4	Dehydrated media are labeled with date of receipt and date opened.
С	12		1.7.5	Caked or expired media or media components are discarded.
С	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.
С	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.
С	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.
С	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.
О	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 Sto	rage 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 I	I - SEAW	ATER	SAMI	PLES

		2.1 Co	llection	and Transportation of Samples
С	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.
С	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.
О	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.
С	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 Ba	cteriolo	gical Examination of Seawater by the APHA MPN
С	9		2.2.1	Lactose broth or lauryl tryptose broth is used as the presumptive medium. (Circle appropriate one.)
С	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 controlNegative 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).
			2.2.6	In a single dilution series, the volumes analyzed are adequate to meet the needs of routine monitoring.
C	6			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.
С	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 Co		l Test for Seawater by APHA MPN
С	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.
		2.3.3	The appropriate positive and negative productivity controls for the presumptive
С	2		media are used. The results are recorded and the records maintained.
C	_	🖵	
		224	Positive productivity controlNegative 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
K	9, 11	🛏	appropriate. (Circle the method of transfer.)
С	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		2.3.9	The presence of turbidity and any amount of gas or effervescence in the culture
C	9		tube constitutes a positive test.
		2.4 Computat	tion of Results – APHA MPN
		2.4.1	Results of multiple dilution tests are read from tables in <i>Recommended</i>
K	9		Procedures for the Examination of Sea Water and Shellfish, Fourth Edition.
		2.4.2	Results from single dilution series are calculated from Hoskins' equation or
K	7		interpolated from Figure 1, Public Health Report 1621 entitled "Most Probable
11	,	-	Numbers for Evaluation of Coli aerogenes Tests by Fermentation Tube
~			Method".
С	7, 9	2.4.3	Results are reported as MPN/100 mL of sample.
		1	ogical Examination of Seawater by the MA-1 Method
С	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-1medium without salicin. Study records are available.
С	5	2.5.3	A-1 medium sterilized for 10 minutes at 121°C.
		2.5.4	The appropriate positive and negative productivity controls for the presumptive
C	2		media are used. The results are recorded and the records maintained.
			Positive productivity controlNegative productivity control
~		2.5.5	Sample and dilutions of sample are shaken vigorously (25 times in a 12" arc in 7
С	9		seconds) before inoculation.
С	9	2.5.6	In a multiple dilution series not less than 3 tubes per dilution are used (5 tubes
	,		are recommended).
С	6	2.5.7	In a single dilution series at least 12 tubes are used.
		2.5.8	In a single dilution series, the volumes analyzed are adequate to meet the needs
			of routine monitoring.
С	6		Sample volume inoculated
		_	Day as of MDN
			Range of MPN
			Strength of media used
		2.5.9	Strength of media used Appropriately diluted process control cultures accompany the samples
C	2		throughout both resuscitation and waterbath incubation Results are recorded
_	_	-	and the records maintained.

			1	
				Positive process control Negative process control
С	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.
С	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 waterbath for the remainder of the 24 \pm 2 hours.
С	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 Cor	nputati	ion 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".
С	7, 9		2.6.3	Results are reported as MPN/100 mL of sample.
				ogical Analysis of Seawater by Membrane Filtration (MF) using
		m'		gar - Materials and Equipment
С	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.
С	23		2.7.2	When using a waterbath for elevated temperature incubation, the level of the water completely covers the plates.
С	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.
С	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 $0.45~\mu m$ and certified by the manufacturer for fecal coliform analyses.
С	2		2.7.7	Lot number, date of receipt and if provided the expiration date of the membrane filters are recorded and records maintained.
С	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.
С	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.
О	11		2.7.12	Forceps tips are clean.
О	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.
С	11		2.7.17	Membrane filter assemblies are autoclave sterilized for 15 minutes at 121°C prior to the start of a filtration series.
О	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 Me	dia Pre	paration and Storage – MF using mTEC Agar
K	11		2.8.1	Phosphate buffered saline is used as the sample diluent and filter funnel rinse.
С	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.
0	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 Sar	nple An	nalyses - MF using mTEC Agar
С	24		2.9.1	mTEC agar is used.
С	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 controlNegative productivity control
С	23		2.9.3	Positive productivity controlNegative productivity control The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration.
C C	23		2.9.3	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before
				The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration.
С	23		2.9.4	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration. The membrane is placed grid side up within the sterile filter apparatus. Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random
C C	23 23, 25		2.9.4	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration. The membrane is placed grid side up within the sterile filter apparatus. 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 C	23 23, 25 23		2.9.4 2.9.5 2.9.6	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration. The membrane is placed grid side up within the sterile filter apparatus. Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling). Sample volumes are filtered under vacuum.
C C K	23 23, 25 23 26		2.9.4 2.9.5 2.9.6 2.9.7	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration. The membrane is placed grid side up within the sterile filter apparatus. Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling). Sample volumes are filtered under vacuum. The pressure of the vacuum pump does not exceed 15 psi. The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration. 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 C K C	23 23, 25 23 26 23, 26		2.9.4 2.9.5 2.9.6 2.9.7 2.9.8	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration. The membrane is placed grid side up within the sterile filter apparatus. Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling). Sample volumes are filtered under vacuum. The pressure of the vacuum pump does not exceed 15 psi. The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration. 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. 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 C K C C	23 23, 25 23 26 23, 26 23		2.9.4 2.9.5 2.9.6 2.9.7 2.9.8 2.9.9	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration. The membrane is placed grid side up within the sterile filter apparatus. Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling). Sample volumes are filtered under vacuum. The pressure of the vacuum pump does not exceed 15 psi. The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration. 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. 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
C C K C C	23 23, 25 23 26 23, 26 23		2.9.4 2.9.5 2.9.6 2.9.7 2.9.8 2.9.9	The sample is shaken vigorously (25 times in a 12" arc in 7 seconds) before filtration. The membrane is placed grid side up within the sterile filter apparatus. Sample volumes tested are consistent with the sampling regime employed (i.e., half log or other appropriate dilutions are used with systematic random sampling). Sample volumes are filtered under vacuum. The pressure of the vacuum pump does not exceed 15 psi. The sides of the filter funnel are rinsed at least twice with 20-30 mL of sterile phosphate buffered saline after sample filtration. 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. 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). Appropriately diluted process control cultures accompany the samples throughout both resuscitation and elevated temperature incubation. Results are

				of resuscitation. Alternatively inoculated plates may be placed in ethafoam prior to air incubation at 44.5 ± 0.5 °C for 24 ± 2 hours.						
			2.9.13	After 2 hours of resuscitation at 35°C, the watertight, tightly sealed containers						
С	11, 23, 24		2.5116	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.						
			2.10.2	Only plates having 80 or fewer colonies are counted. If it is unavoidable to use						
С	23	Ш		plates having more than 80 colonies, counts are given as >80 x 100/the volume						
		 	2.10.3	of sample filtered. When multiple dilutions are filtered, the laboratory has developed a procedure						
C	2, 11, 23		2.10.3	for assessing the contribution of all positive dilutions to the final count.						
			2.10.4	The number of fecal coliforms is calculated by the following equation:						
a	22 11									
С	23, 11	Ш		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.						
С	23, 11		2.10.5	Results are reported as CFU/100 mL of sample.						
PART I	II - SHEL	LFISI	H SAM	PLES						
		3.1 Co	llection	and Transportation of Samples						
С	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						
K	,			containers loosely sealed.						
*77			3.1.3	Shellstock samples are labeled with collector's name, type of shellstock, the						
K	9	ш		source or harvest area, sampling station, time, date and place (if applicable) of collection.						
			3.1.4	Immediately after collection, shellfish samples are placed in dry storage (ice						
a			3.1.1	chest or equivalent) which is maintained between 0 and 10°C with ice or cold						
С	9	Ш		packs for transport to the laboratory. Once received, the samples are placed						
				under refrigeration unless processed immediately.						
~			3.1.5	Analysis of the samples is initiated as soon as possible after collection. Shellfish						
С	1	ш		samples are not tested if the time interval between collection and analysis exceeds 24 hours.						
		2 2 Dw	norotio	on of Shellfish for Examination						
		3.2 FT	3.2.1							
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.						
			3.2.3	The hands of the analyst are thoroughly washed with soap and water						
О	9			immediately prior to cleaning the shells of debris.						
О	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						
			0.0.1	drinking water quality.						
О	9		3.2.6	Shellstock are allowed to drain in a clean container or on clean towels prior to						
			3.2.7	opening. Immediately prior to shucking, the hands (or gloved hands) of the analyst are						
K	9		3.2.7	thoroughly washed with soap and water and rinsed in 70% alcohol.						
С	9	П	3.2.8	Shellstock are not shucked directly through the hinge.						
_			3.2.9	Contents of shellstock (liquor and meat) are shucked into a sterile, tared blender						
С	9			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	П	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
K	2			diluent is added.
О	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 Recommended Procedures for the Examination of Sea Water And Shellfish, Fourth Edition is followed for the analysis of previously shucked and frozen shellfish meats.
	1	3.3 MI	N Anal	ysis for Fecal Coliform Organisms, Presumptive Test, APHA
		<u> </u>	3.3.1	Appropriate strength lactose or lauryl tryptose broth is used as presumptive
С	9		3.3.1	media in the analysis. (Circle the medium used.)
С	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 controlNegative 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.
С	9		3.3.4	No fewer than 5 tubes per dilution are used in a multiple dilution MPN series.
С	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
С	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 Co	nfirmed	Test for Fecal Coliforms - APHA
C	9		3.4.1	EC medium is used as the confirmatory medium.
С	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.
			0.40	Positive productivity controlNegative 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.)
С	9		3.4.4	EC tubes are incubated in a circulating waterbath at 44.5 ± 0.2 °C
K	9		3.4.5	EC tubes are read for gas production after 24 ± 2 hours of incubation.
С	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 Co	mputati	on 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".
С	9		3.5.3	Results are reported as MPN/100 grams of sample.
		3.6 Sta	ndard I	Plate Count Method
О	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.
С	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.
С	9		3.6.7	Not more than 1 mL nor less than $0.1\ \text{mL}$ 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 Co		on 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 Bac	cteriolo	gical 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.
С	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.

С	2, 3		3.8.6	The phosphate buffered saline is tempered at 45 - 50°C to prevent premature solidification of the agar.
С	9			
			3.8.8	Six grams of shellfish (12 grams of homogenate if initially diluted 1:1) is placed
С	2,3		2.0.0	into a sterile container and the contents brought up to 60 mL with sterile,
	,	_		tempered phosphate buffered saline.
17	2		3.8.9	Sixty (60) mL of tempered, twice boiled double strength Modified MacConkey
K	3			Agar is added.
K	2,3, 22		3.8.10	The container is gently swirled or slowly inverted once to mix the contents,
IX	2,3, 22			which are subsequently distributed uniformly over six plates.
С	1		3.8.11	Media and diluent sterility are determined with each use. Results are recorded
-	_	<u> </u>		and the records maintained.
			3.8.12	Media productivity is determined using media appropriate properly diluted
a	1			pour plated positive and negative control cultures for each batch of Modified
С	1	ш		MacConkey agar prepared.
				Positive control culture Negative control culture
			3.8.13	When solidified, the plates are placed inverted into an air incubator at $45.5 \pm$
С	3, 13	ш	3.0.13	0.5°C for 18 to 30 hours of incubation.
С	2		3.8.14	Plates are stacked no more than three high in the incubator.
-			3.8.15	Appropriately diluted pour plated process control cultures accompany each set
				of samples throughout incubation. The results are recorded and the records
C	2			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
11	11			magnification and visibility for counting.
О	1		3.9.2	A hand tally or its equivalent is used to aid in counting.
С	3, 6		3.9.3	All brick red colonies greater than 0.5 mm in diameter are totaled over all the
	3,0			plates and multiplied by a factor of 16.7.
C	3		3.9.4	Results are reported as CFU/100 grams of sample.
		Bacter	iologica	l Examination of Soft-shelled Clams and American Oysters for Male
			Specific	Coliphage (MSC)
		3.10 M	SC Equ	ipment and Supplies
K	30	1		Sample containers used for the shucked sample are sterile, made of glass or
K	30	Ш		some other inert material (i.e. polypropylene) and hold $100 - 125$ mL.
			3.10.2	The refrigerated centrifuge used must have the capacity to accommodate the
C	27, 28			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
			0.10.4	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
	<u> </u>		2 10 5	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.
	<u> </u>	<u> </u>	3.10.6	The sterility of each batch of reusable glass syringes is determined. Results are
K	1		5.10.0	recorded and records maintained.
		_	1	

			_	
С	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.
С	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 M	SC Med	lia 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.
С	27, 28		3.11.4	The streptomycin and ampicillin solutions are added to tempered bottom agar and vortex for 2 minutes on stir plate.
О	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 Pr	eparati	on 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.
О	2		3.12.2	The blades of shucking knives are not corroded.
О	9		3.12.3	The hands of the analyst are thoroughly washed with soap and water immediately prior to cleaning the shells of debris.
О	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.
О	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.
С	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			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 M	ISC Sa	mple Analysis
С	28		3.13.1	E.coli <i>Famp</i> ATCC 700891 is the bacterial host strain used in this procedure.
K	27, 28		3.13.2	Host cell growth broth is tempered at $36 \pm 1^{\circ}$ 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 $36 \pm 1^{\circ}$ C for 4-6 hours to provide host cells in log phase growth for sample analysis.
С	27, 28		3.13.4	After inoculation, the host cell growth broth culture is not shaken.
С	28		3.13.5	A 2:1 mixture of sterile growth broth to shellfish tissue is used for eluting the MSC.

С	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.
С	28		3.13.7	The elution mixture is homogenized at high speed for 180 seconds.
С	28		3.13.8	Immediately after blending, 33 grams of the homogenized elution mixture are weighed into centrifuge tubes.
С	28		3.13.9	The homogenized elution mixture is centrifuged for 15 minutes at 9000 x g at 4°C.
C	27, 28		3.13.10	The supernatant is pipetted off, weighed and the weight recorded.
С	27, 28		3.13.11	The supernatant is allowed to warm to room temperature about $20\ \mathrm{to}\ 30\ \mathrm{minutes}.$
K	27, 28		3.13.12	The autoclaved soft agar is tempered and held at 51 \pm $1^{\circ}C$ throughout the period of sample analysis.
K	27, 28		3.13.13	Two hundred microliters (0.2 mL) of log phase host strain $E\ coli$ 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.
С	27, 28		3.13.15	2.5 mL of sample supernatant is added to each tube of tempering soft agar.
С	27, 28		3.13.16	The soft agar/sample supernatant/host cell mixture is gently rolled between the palms of the hands to mix.
С	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.
С	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.
K	27, 28		3 13 20	Positive control Growth broth is used as the negative control or blank.
K	21, 28		+	Type strain MS2 (ATCC 15597) male specific bacteriophage appropriately
K	27, 28			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.
С	27, 28		3.13.24	All plates are incubated at $36 \pm 1^{\circ}$ C for 18 ± 2 hours.
		3.14 C	omputat	ion of Results - MSC
С	27		3.14.1	Circular zones of clearing or plaques of any diameter in the lawn of host bacteria are counted.
С	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.
О	9		3.14.4	The MSC count is rounded off conventionally to give a whole number.

REFERENCES

- 1. American Public Health Association 1984. *Compendium of Methods for the Microbiological Examination of Foods*, 2nd Edition. APHA, Washington, D.C.
- 2. Good Laboratory Practice.
- 3. "Interim Guides for the Depuration of the Northern Quahog, *Mercenaria mercenaria*." 1968. Northeast Marine Health Sciences Laboratory, North Kingstown, RI.
- 4. U.S. Department of Commerce. 1976. NBS *Monograph 150*. U.S. Department of Commerce, Washington, D.C.
- Association of Official Analytical Chemists (AOAC). Current Edition. Official Methods of Analyses of the Association of Official Analytical Chemists. Official method 978.23. Chapter 17.305. AOAC Arlington, VA
- 6. Wilt, D.S. (ed.). 1974. *Proceedings of the 8th National Shellfish Sanitation Workshop*. U.S. Food and Drug Administration, Washington, D.C.
- 7. U.S. Public Health Service (PHS). 1947. Public Health Report, Reprint #1621. PHS, Washington, D.C.
- 8. Association of Official Analytical Chemists (AOAC). 1991. *Quality Assurance Principles for Analytical Laboratories*. AOAC, Arlington, VA.
- 9. American Public Health Association (APHA). 1970. *Recommended Procedures for the Examination of Sea Water and Shellfish*, 4th Edition. APHA, Washington, D.C.
- 10. Interstate Shellfish Sanitation Conference (ISSC). 1986. *Shellfish Sanitation Interpretation #SS-39*. ISSC, Columbia, S.C.
- 11. American Public Health Association (APHA). 1992. *Standard Methods for the Examination of Water and Wastewater*, 18th Edition. APHA/AWWA/WEF, Washington, D.C.
- 12. Title 21, Code of Federal Regulations, Part 58, *Good Laboratory Practice for Nonclinical Laboratory Study*. U.S. Government Printing, Washington, D.C.
- 13. American Public Health Association (APHA). 1992. *Standard Methods for the Examination of Dairy Products*, 16th Edition. APHA, Washington, D.C.
- 14. Fisher, J. 1985. Measurement of pH. American Laboratory 16:54-60.
- 15. Consult pH electrode product literature.
- 16. Association of Official Analytical Chemists (AOAC). 1999. AOAC Methods Validation and Technical Programs Criteria for Laboratories Performing Food Testing. AOAC, Arlington, VA.
- 17. U.S. Environmental Protection Agency (EPA). 1975. *Handbook for Evaluating Water Bacteriological Laboratories*. EPA-670/9-75-006. U.S. EPA, Cincinnati, OH
- 18. Adams, W.N. 1974. NETSU. Personal communication to Dr. Wallace Andrews, FDA.
- 19. U.S. Food and Drug Administration (FDA).1995.*Bacteriological Analytical Manual*. U.S. FDA, 8th Edition, AOAC, Arlington, VA.
- 20. U.S. Food and Drug Administration (FDA) and Interstate Shellfish Sanitation Conference (ISSC). 1997. *NSSP Guide to the Control of Molluscan Shellfish*. FDA/ISSC, Washington, D.C. and Columbia, S.C.
- 21. U.S. Environmental Protection Agency. 1978. *Microbiological Methods for Monitoring the Environment, Water and Wastes*. EPA/600/8/78/017. EPA, Washington, D.C.
- 22. Furfari, Santo. March 21, 1972. Personal Communication to Dan Hunt, FDA.
- 23. United States Environmental Protection Agency, *Improved Enumeration Methods for the Recreational Water Quality Indicators: Enterococci and Escherichia coli*. EPA/821/R-97-004, EPA, Washington, DC
- 24. Rippey, Scott, R, Adams, Willard, N, and Watkins, William, D. Enumeration of fecal coliforms and *E. coli* in marine and estuarine waters: an alternative to the APHA-MPN approach, Journal WPCF, 59, 8 (1987).
- 25. FDA Manual of Interpretations, National Shellfish Sanitation Program *Guide for the Control of Molluscan Shellfish*, 2003 Revision, Interpretation Number 03-IV-@.02-102.
- 26. Membrane filtration: A Users Guide and Reference Manual, Thomas D. Brock, Science Tech Inc., Madison, WI, 1983.
- 27. Proceedings of the Male-specific Bacteriophage (MSC) Workshop, Gloucester, MA, March 9-12, 2004.

- 28. MSC Method and SLV write-up, Proposal 05-114 Spinney Creek Shellfish, Inc., September, 2009.
- 29. American Public Health Association. 1970. *Recommended Procedures for the Examination of Sea Water and Shellfish*, 4th Edition, APHA, New York, N.Y.
- 30. ASTM Manual on the Use of Thermocouples in Temperature Measurement, MNL-12 (ASTM, West Conshohocken, PA, 1993).
- 31. JOHN KAROLUS, MERCURIA CUMBO, SUSAN BOEHLER, and LAURA SAVINA. Modification of an Approved Medium for Fecal Coliform Detection in Seawater: A-1 Medium Minus Salicin. *Journal of Food Protection*: Vol. 66, No. 1, pp. 120–121.
- 32. MSC Method and SLV write-up, Proposal 13-120 Spinney Creek Shellfish, Inc., January, 2014.

	ARY OF NONC	CONFORMITIES	
age	Item	Observation	Documentation Required
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			i

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 Micro	biological Component:
1. Does Not Conform Status : The Microbiological componen requirements if:	t of this laboratory is not in conformity with NSSP
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	
2. Provisionally Conforms Status : The microbiological comp	onent of this
laboratory is determined to be provisionally conforming to NS	SP requirements if
the number of critical nonconformities is ≥ 1 but ≤ 3	
C. Laboratory Status (<i>circle appropriate</i>)	
ci Zucoraiory Zuitas (en ete appropriate)	
Does Not Conform Provisionally Conforms Confor	ms
Acknowledgment by Laboratory Director/Supervisor:	
All corrective Action will be implemented and verifying substated Evaluation Officer on or before	ntiating documentation received by the Laboratory
Laboratory Signature: Date:	
LEO Signature: Date:	

NSSP Form LAB-100 Microbiology Rev. 2010-11-08

Laboratory Evaluation Checklist - PSP

PUBLIC HEALTH SERVICE				
J.S. FOOD AND DRUG ADMINISTRATION				
SHELLFISH PROGRAM IMPLEMENTAT	ION BRANCH			
SHELLFISH SAFETY TEAM				
5100 PAINT BRANCH PARKWAY				
COLLEGE PARK, MD 20740-3835				
ΓΕL. 301-436-2151/2147 FAX 301-436-2672				
SHELLFISH LABORATORY EVALUATIO	ON CHECKLIST			
LABORATORY:				
ADDRESS:				
TELEPHONE: FAX:	EMAIL:			
DATE OF EVALUATION:	DATE OF REPORT:	LAST EVALUATION:		
LABORATORY REPRESENTED BY:	TITLE:			
LABORATORY EVALUATION OFFICER:	SHELLFISH SPECIA	ALIST:		
	REGION:			
OTHER OFFICIALS PRESENT:	TITLE:			
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(Assessment of the second of t				
tems which do not conform are noted by:				
C- Critical K - Key O - Other NA - Not Applica	shla Canformity is mater	1 by a "12"		
C- Chucai K - Key O - Other NA - Not Applica	able Comornity is noted	ı uy a v		

	PART I - QUALITY ASSURANCE						
Code	-						
	Quality Assurance (QA) Plan						
K	1. Written Plan adequately covers all the following: (check √ those that apply)						
	1. a. Organization of the laboratory.						
	2. b. Staff training requirements.						
	3. c. Standard operating procedures.						
	4. d. Internal quality control measures for equipment, calibration, maintenance, repair and						
	performance.						
	5. e. Laboratory safety.						
	6. f. Quality assessment.						
	7. g. Proper animal care.						
$\overline{\mathbf{C}}$	2. QA plan implemented.						
	1.2 Work Area						
О	1. Adequate for workload and storage.						
О	2. Clean and well lighted.						
О	3. Adequate temperature control.						
О	4. All work surfaces are nonporous and easily cleaned.						
C	5. A separate, quiet area with adequate temperature control for mice acclimation and injection						
	is maintained.						
	1.3 Laboratory Equipment						
O	1. The pH meter has a standard accuracy of 0.1 unit.						
K	2. pH paper in the appropriate range (i.e. 1-4) is used with minimum accuracy of 0.5 pH units.						
K	3. pH electrodes consist of pH half cell and reference half cell or equivalent combination electrode						
	(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	4. pH meter is calibrated daily or with each use. Records maintained.						
K	5. Effect of temperature has been compensated for by an ATC probe or by manual adjustment.						
K	6. A minimum of two standard buffer solutions (2 & 7) is used to calibrate the pH meter. Standard						
	buffer solutions are used once and discarded.						
K	7. Electrode efficiency is determined daily or with each use following either slope or millivolt						
TZ	procedure.						
K	8. The balance provides a sensitivity of at least 0.1g at a load of 150 grams.						
K	9. The balance calibration is checked monthly using NIST Class S or ASTM Class 1 or 2 weights or						
IZ	equivalent. Records maintained.						
K	10. Refrigerator temperature is maintained between 0 and 4°C.						
0	11. Refrigerator temperature is monitored at least once daily. Record maintained.						
K	12. Freezer temperature is maintained at -20°C or below.						
0	13. Freezer temperature is monitored at least once daily. Record maintained.						
0	14. All glassware is clean.						
О	15. Once during each day of washing, several pieces of glassware from each batch washed are tested						
	for residual detergent with aqueous 0.04% bromthymol blue solution. Records are maintained.						

	1.4 Reagent and Reference Solution Preparation and Storage
C	1. Opened PSP reference stand solution (100 μg/ml) is not stored.
K	2. PSP working standard solution (1 μg/ml) and all dilutions are prepared with dilute HCl, pH 3 water, using 'Class A' volumetric glassware (flasks and pipettes) or prepared gravimetrically.
K	3. Refrigerated storage of PSP working standard solution (1µg/ml) does not exceed 6 months and is checked gravimetrically for evaporation loss.
K	4. PSP working dilutions are discarded after use.
K	5. Make up water is distilled or deionized (<i>circle one</i>) and exceeds 0.5 megohm resistance or is less than 2 μ Siemens/cm conductivity at 25°C to be tested and recorded monthly for resistance or conductivity (<i>circle the appropriate</i>).
0	6. Make up water is analyzed for residual chlorine monthly and is at a nondetectable level (≤ 0.1 ppm). Records maintained.
K	7. Make up water is free from trace (< 0.5 mg/l) dissolved metals specifically Cd, Cr, Cu, Ni, Pb, and Zn as determined annually with total heavy metal content \leq 1.0 mg/l. Records maintained.
О	8. Makeup water contains < 1000 CFU/ml as determined monthly using the heterotrophic plate count method. Records maintained
	1.5 Collection and Transportation of Samples
O	1. Shellstock are collected in clean, waterproof, puncture resistant containers.
K	2. Samples are appropriately labeled with the collector's name, harvest area and time and date of collection.
K	3. Immediately after collection, shellstock samples are placed in dry storage for transport (e.g. cooler) which is maintained between 0 and 10°C. Upon receipt at the lab, samples are placed under refrigeration.
K	4. The time from collection to completion of the bioassay should not exceed 24 hours. However, if there are significant transportation delays, then shellstock samples are processed immediately as follows (<i>circle the appropriate choice</i>):
	a. Washed, shucked, drained, frozen until extracted;
	b. Washed, shucked, drained, homogenized and frozen;
	c. Washed, shucked, drained, extracted, the supernatant decanted and refrigerated (best choice); or
	d. The laboratory has an appropriate contingency plan in place to handle samples which can't be analyzed within 24 hours due to transportation issues.
K	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.
PAF	RT II - EXAMINATION OF SHELLFISH FOR PSP TOXIN
	2.1 Preparation of Sample
C	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	2. The outside of the shell is thoroughly cleaned with fresh water.
	3. Shellstock are opened by cutting adductor muscles.
	4. The inside of the shell is rinsed with fresh water to remove sand or other foreign material.
0	5. Shellfish meats are removed from the shell by separating adductor muscles and tissue connecting at
	p. shemish means are removed from the shell by separating addition inductes and tissue connecting at

		the hinge.
K		6. Damage to the body of the mollusk is minimized in the process of opening.
O		7. Shucked shellfish are drained on a #10 mesh sieve (or equivalent) without layering for 5 minutes.
K		8. Pieces of shell and drainage are discarded.
$\overline{\mathbf{C}}$		9. Drained meats or thawed homogenates are blended at high speed until homogenous (60 - 120
		seconds).
		2.2 Extraction
K		1. 100 grams of homogenized sample is weighed into a beaker.
K		2. An equal amount of 0.1 N/0.18 N HCl is added to the homogenate and thoroughly mixed (circle the
		appropriate normality).
C		3. pH is checked and, if necessary adjusted to between pH 2.0 and 4.0.
C		4. Adjustment of pH is made by the dropwise addition of either the acid (5 N HCl) or base (0.1N
		NaOH) while constantly stirring the mixture.
C		5. The homogenate/acid mixture is promptly brought to a boil, $100 \pm 1^{\circ}$ C, then gently boiled for 5
		minutes.
0		6. The homogenate/acid mixture is boiled under adequate ventilation (i.e. fume hood).
O		7. The extract is cooled to room temperature.
C	L	8. The pH of the extract is determined and adjusted, if necessary to between pH 2 and 4,
		preferably to pH 3 with the stirred dropwise addition of 5 N HCl to lower the pH or 0.1N NaOH
TZ		to raise the pH.
K		9. The extract volume (or mass) is adjusted to 200 mls (or grams) with dilute HCl, pH 3 water.
K	_	10. The extract is returned to the beaker, stirred to homogeneity and allowed to settle to remove particulates; or, if necessary, an aliquot of the stirred supernatant is centrifuged at 3,000 RPM for 5
		minutes before injection.
K		11. If mice cannot be injected immediately then the supernatant should be removed from the centrifuge
1.		tubes and refrigerated for up to 24 hours.
K		12. Refrigerated extracts are allowed to reach ambient temperature before being bioassayed.
		2.3 Bioassay
O	Г	1. A 26-gauge hypodermic needle is used for injection.
K		2. Healthy mice in the weight range of 17 -23 grams (19 - 21 grams preferable) from a stock colony are
		used for routine assays. Mice are not reused for bioassay.
		Stock strain usedSource of mice
$\overline{\mathbf{C}}$		3. Mice are allowed to acclimate for at least 24 hours prior to injection. In some cases up to 48
		hours may be required.
C		4. A conversion factor (CF) has been determined as Month and year when current
		CF determined
C	L	5. CF value is checked weekly if assays are done on several days during the week, or, once each
		day that assays are performed if they are performed less than once per week.
		Date of most recent CF check
		CE varified/CE not varified (Civale appropriate alreion)
		CF verified/CF not verified (Circle appropriate choice) (If the CF is not verified 5 additional miss are injected with the dilution used in the CF sheet to
C		6. If the CF is not verified, 5 additional mice are injected with the dilution used in the CF check to complete a group of 10 mice. Ten additional mice are also injected with this dilution to produce a
1		rompiem a group of to mice. Ten auditional mice are also injected with tins undutin to produce a

	3 44 4 77 6774 1 1 1 4 1 4 1 4 1
	second group of 10 mice. The CF is calculated for each group of 10 mice and averaged to give the CF to be used in sample toxicity calculations for the day's or week's work only. All subsequent
	work must make use of the original laboratory CF value unless this value continues to fail to be
	verified by routine CF checks.
C	7. If the CF fails to be verified, the cause is investigated and the situation corrected. If the cause
	cannot be determined with reasonable certainty and fails > 3 times per year, the bioassay is
	restandardized.
O	8. Mice are weighed to the nearest 0.5 gram.
C	9. Mice are injected intrapertioneally with 1 ml of the acid extract.
K	10. For the CF check, at least 5 mice are used.
C	11. At least 3 mice are used per sample in routine assays.
C	12. Elapsed time is accurately determined and recorded.
K	13. If death occurs, the time of death to the nearest second is noted by the last gasping breath.
C	14. If median death time(2 out of 3 mice injected die) is < 5 minutes, a dilution is made with dilute
	HCl, pH 3 water, to obtain a median death time in the range of 5 to 7 minutes.
	2.4 Calculation of Toxicity
C	1. The death time of each mouse is converted to mouse units (MU) using Sommer's Table (Table 6
	Recommended Procedures, 4 th edition). The death time of mice surviving beyond 60 minutes is
	considered to be < 0.875 MU.
K	2. A weight correction in MU is made for each mouse injected using Table 7 in <i>Recommended</i>
	Procedures, 4 th edition.
C	3. The death time of each mouse in MU is multiplied by a weight correction in MU to give the
	corrected mouse unit (CMU) for each mouse.
C	4. The median value of the array of corrected mouse units (CMU) is determined to give the
	median corrected mouse unit (MCMU).
C	5. The concentration of toxin is determined by the formula, MCMU x CF X Dilution Factor X
	200.
C	6. Any value greater than 80µg/100 grams of meat is actionable.

REFERENCES

- 1. Adams, W.N. and S.A. Furfari. 1984. Evaluation of laboratory performance of the AOAC method for PSP toxin in shellfish. *J. Assoc. Off. Anal. Chem.* Vol 67, 6:1147-1148.
- 2. American Public Health Association. 1970. *Recommended Procedures for the Examination of Sea Water and Shellfish*, 4th Edition. APHA, Washington, D.C.
- 3. American Public Health Association. 1992. *Standard Method for the Examination of Dairy Products*, 16th Edition. APHA, Washington, D.C.
- Association of Official Analytical Chemists International. 1990. *Methods of Analysis*, 15th Edition. AOAC, Arlington, VA.
- 5. APHA/WEF/AWWA. 1992. Standard Methods for the Examination of Water and Wastewater, 18th Edition. APHA, Washington, D.C.
- 6. Title 21, Code of Federal Regulations, Part 58, *Good Laboratory Practice for Nonclinical Laboratory Study*. U.S. Government Printing, Washington, D.C.
- 7. National Research Council. 1996. *Guide for the Care and Use of Laboratory Animals*. National Academy Press, Washington, D.C.

National Shellfish Sanitation Program (NSSP) Guide for the Control of Molluscan Shellfish: 2013 Revision				
8.	Personal communication with USFDA Washington Seafood Laboratory Branch, Office of Seafood, CFSAN, 1998-1999.			

LABORATORY:			D	DATE OF EVALUATION:		
SHELLFISH LABORATORY EVALUATION CHECKLIST						
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Page	Item	Observation		Documentation Required		
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PARALYTIC SHELLFISH POISON COMPONENT: PA	RTS 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 PS	SP Component			
1. Does Not Conform Status The PSP component of this labor requirements if:	oratory is not in conformity with NSSP			
A. The total # of Critical nonconformities is ≥ 3 or				
B. The total # of Key nonconformities is ≥ 6 or				
C. The total # of Critical, Key and Other is ≥ 10				
2. Provisionally Conforms Status : The PSP component of this laboratory is determined to be provisionally conforming to NSSP requirements if the number of critical nonconformities is ≥ 1 but < 3				
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 Rev. 2005-08-19

Laboratory Evaluation Checklist - Analysis for NSP (Mouse Bioassay)

PUBLIC H	PUBLIC HEALTH SERVICE					
U.S. FOOD	AND DRUG	ADMIN	ISTRATI	(ON		
				TION BRANCH		
	H SAFETY					
	T BRANCH					
	PARK, MD			_		
	36-2151/2147					
		TORY E	VALUAT	ION CHECKLIST		
LABORAT						
ADDRESS:						
TELEPHO	NE:	FA	X:	EMAIL:		
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Items which	Items which do not conform are noted by:					
	Total to all company and all all all all all all all all all al					
C- Critical K	C- Critical K - Key O - Other NA- Not Applicable Conformity is noted by a " $$ "					
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Weighted Code	Item Description		
	Quality Assurance (QA) Plan		
С	 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. External performance assessment. 		
С	2. QA Plan is implemented		
	Work Area		
О	1. Adequate for workload and storage.		
O	2. Clean and well lighted.		
О	3. All work surfaces are nonporous and easily cleaned.		
С	4. A separate, quiet area with adequate temperature control is maintained for acclimation and injection of mice.		
	Laboratory Equipment		
K	 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 used. 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 sensitivity of 0.1 g at a load of 20 g is required. 		
O	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.		
О	2. Reagent grade NaCl is used in the extraction procedure.		
С	3. Diethyl ether purified for lipid extraction is used for extracting lipids from the shellfish homogenates.		
С	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
O	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 the time and date of collection.
K	3. Immediately after collection, shellstock samples are placed in dry storage between 0 and 10°C until analyzed.
K	4. Shellstock samples are analyzed within 24 hours of collection or refrigerated unshucked until analyzed.
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 prepared according to steps 1-9 in "Preparation of Sample" section below.
	Preparation of Sample
$\overline{\mathbf{C}}$	1. At least 12 animals are used per sample.
O	2. The outside of the shell is thoroughly cleaned with fresh water.
K	3. Shellstock are opened by cutting the adductor muscles.
$\overline{\mathbf{C}}$	4. Shell liquor is discarded.
O	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.
O	9. Shucked shellfish are drained on a #10 mesh sieve or equivalent without layering for 5 minutes.
K	10. Pieces of shell and drainings are discarded.
$\overline{\mathbf{C}}$	11. Drained meats are blended at high speed until homogenous (60-120 seconds).
$\overline{\mathbf{C}}$	12. Shellfish homogenates are digested within 2 hours of blending.
	Digestion 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.
С	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 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 reached, gently boil for 5 minutes.
О	5. The beaker is covered with a watch glass or equivalent during boiling to prevent excessive evaporation.
O	6. The homogenate is boiled under adequate ventilation (fume hood).
O	7. The boiled, acidified homogenate is cooled to room temperature or below in a refrigerator or in an ice bath.
	Extraction
C	1. All steps in the extraction procedure which involve any manipulation of diethyl ether are carried out under adequate ventilation.

C	2. 100 ml of diethyl ether is added to the cooled, acidified homogenate in a stoppered centrifuge tube and shaken vigorously for 5 minutes.
O	3. Centrifuge tubes are vented frequently while being shaken and before being centrifuged to avoid accidents.
C	4. The content of the centrifuge tubes are centrifuged at 2000 rpm for 10 to 15 minutes.
$\overline{\mathbf{C}}$	5. The clear upper ether phase is transferred to a large separatory funnel.
С	6. The contents of the centrifuge tube are extracted three additional times for 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.
	Bioassay
С	1. The volume of the lipid residue is adjusted by weight to 10 ml (9.17 g) per 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
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
O	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.
	8. A total of 5 mice are injected with undiluted or diluted extract as appropriate per sample in routine assays.
С	 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 analysis.
C	9. The time of completed injection is recorded.
C	10. 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 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 continuous observation.
	Calculation of Toxicity
C	1. The death time of each mouse is converted to mouse units (MU) using Table 8 in <i>Recommended Procedures</i> , 4 th Edition.
O	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 <i>Recommended Procedures</i> , 4 th Edition.
O	4. Table 8 is interpolated to accommodate weights which are not listed.
C	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:		
SHELLFISH LABORATORY EVALUATION CHECKLIST SUMMARY OF NONCONFORMITIES					
Page	Item	Observation	Documentation Required		
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LABORATORY STATUS					
LABORATORY	DA	ATE			
LABORATORY REPRESENTATIVE:					
NEUROTOXIC SHELLFISH POISON COMPO	NENT:				
A. Results					
Total # of Critical (C) Nonconformities					
Total # of Key (K) Nonconformities	_				
Total # of Critical, Key and Other (O) nonconformit					
B. Criteria for Determining Laboratory Status	s of the NSP Component				
1. Does Not Conform Status The NSP component of this laboratory is not in conformity with NSSP requirements if:					
A. The total # of Critical nonconformities is ≥ 3 or					
B. The total # of Key nonconformities is ≥ 6 or					
C. The total # of Critical, Key and Other is ≥ 10					
2. 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 < 3					
C. Laboratory Status (circle appropriate)					
Does Not Conform Provisionally Conform	s Conforms				
Acknowledgment by Laboratory Director/Supervisor	r:				
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 Analysis for NSP (Mouse Bioassay) 2005-08-19