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

**COLLEGE PARK, MD 20740-3835** TEL. 240- 402-2151/2055/4960 FAX 301-436-2601

CFSANDSSLEOS@FDA.HHS.GOV						
	SHELLFISH LA	ABORATOR	RY EVA	LUATION	CHECKLIST	
LABORA'	ΓORY:					
ADDRESS	<b>5</b> :					
TELEPHO	ONE:		FAX:			
EMAIL:			•			
DATE OF	EVALUATION:	DATE OF	REPOI	RT:	LAST EVALUATION	<b>N</b> :
LARORA'	TORY REPRESENTED	) RV·	ŀ	TITLE:		
<u> </u>	TORT REPRESENTED	, , , , , , , , , , , , , , , , , , , ,		11122,		
LABORA'	TORY EVALUATION	OFFICER:	À	SHELLFISH	I SPECIALIST:	
OTHER C	OFFICIALS PRESENT:		r	TITLE:		
OTTILITY	TITOTIES TRESELLE			11122,		
that for all separate re summary	ty is noted by a (Y), no (I I N/A indications, you m ecord. Record comments of nonconformities. All n ast be in place for onsite	ust documen s related to a nonconformi laboratory e	nt the ro any req ities mu	eason why th uirement on ast be identif ion to be sch	nis requirement is N/A the space provided in ied and explained. Qua	on a the
		Parts of the				
Part I	Quality Management: La					
Part II	Quality Assurance: The I					m
Part III	I Quality Control: Documentation for Quality System Defensibility					

_	-	anagement: Laboratory Operations and Responsibilities for nitation Program Laboratory Quality Systems
		ITEM
Conformance Comments	Ref	
		1.1 Components of the Laboratory Quality System
	1,3,6,9	1.1.1 The laboratory has an overall Quality System supported by quality management structure, quality assurance processes and quality control functions.
	1,3,6,9	1.1.2 Management and technical structure exist to support the Quality System.
	1,3,6,9	1.1.3 Quality documentation is required by the laboratory. These include a Quality Assurance (QA) Manual (or otherwise named) and Standard Operating Procedures (SOPs) to support the quality assurance process of the laboratory.
	1,9	1.1.4 The <u>documents</u> used to implement the quality assurance process and <u>records</u> used to verify quality control (QC) function of the laboratory are reviewed and controlled.
	9	1.1.5 An established process of Quality System assessment and technical proficiency are documented with results retained until the next review.
	9	1.1.6 Resolution, management review and prevention of nonconformities are a documented component of the Quality System.
		1.2 Laboratory Management Structure and Quality Systems
	1,3,6,9	1.2.1 The laboratory's structure is clearly organized with supervisory chain delineated.
	9	1.2.2 The laboratory has ensured that its management and personnel are free from any undue internal and external commercial, financial and other pressures and influences that may adversely affect the quality of their work.
	9	1.2.3 The laboratory has documentation of dedicated personnel with the authority and resources required to carry out their duties, including implementing and maintaining the Quality System of the laboratory.
	1, 9	1.2.4 The laboratory's designated quality personnel ensure adherence to the quality system, including SOPs and QC. These staff have clear documented authority to initiate actions to prevent or minimize departures from quality system and monitor the corrective action process.
	9	1.2.5 The laboratory has documentation of a designated quality system manager, responsible for monitoring all aspects of the quality system to assure defensibility. This person shall have unrestricted access to FDA Shellfish Laboratory Evaluation Officers (LEOs) and the highest levels of the laboratories management. In the case of a single person laboratory, FDA
	1, 9	LEOs will assist with developing a monitoring plan.  1.2.6 A documented system is in place to ensure that appropriate
	1, 9	review of and communication regarding the elements of the quality system are established among the laboratory staff and laboratory management.
		1.3 Laboratory Personnel and Roles in a Quality System
	1,3, 9	1.3.1 The roles and responsibilities of all personnel are defined in the QA

			manual, read by all staff and the acknowledgments of these
			responsibilities are retained.
	9	1.3.2	The laboratory policy and the training procedures for personnel are documented and relevant to the scope of the current activities in the laboratory. If the laboratory intends to add methods to their scope, training SOPs must also be added with successful completion by the analyst(s) that will perform the method(s). In the case of a single person laboratory, method proficiency verification must be retained during the life of the methods use in the laboratory.
	9	1.3.3	The laboratory shall maintain a personnel file/ record of any relevant authorization(s), qualifications, trainings, and/or proficiencies for each analyst. This information shall be available upon request as verification of staff training and shall be retained for all staff until two years after they are no longer employed by the laboratory.
	1, 3, 9	1.3.4	The laboratory has documented that all personnel involved in testing have read and understand the applicable SOPs and associated quality documentation and implement the policies and procedures required for the performance of their technical function.
PART II – (	Quality Ass	urance: '	The Process of Documenting and Maintaining a Quality System
			ality Assurance Process: QA Manual, SOPs and Document
	1.0	Contro	
	1, 9	2.1.1	The QA manual shall include or make reference to all laboratory SOPs and any supporting procedures, including technical procedures.
	1, 9	2.1.2	SOPs are controlled documents and include detailed, written instructions to achieve uniformity of test methods and quality control procedures, such that items that might affect the quality of the outcome are mitigated.
	1, 9	2.1.3	SOPs and the QA Manual are controlled documents, such that specific individuals are designated within the laboratory with editorial control. These individuals are identified in the QA Manual.
	1,9	2.1.4	Each time an SOP or the QA manual has changed, the new version will be marked as such and will be distributed to the laboratory with older versions removed from circulation.
	1, 9	2.1.5	Staff training requirements are documented in the QA manual and
		220	the training procedure is included.
	1,9	2.2 Qua 2.2.1	Ality Manual Items  Quality Assurance Manual contains:  Table of Contents;  Organizational chart;
			A description of the Quality System and procedure for implementation and maintenance; Policy and procedure for resource management (human resources,
			competence and training, work environment and safety), description of responsibilities;
			Policy and procedures for rejection criteria; Policy and procedures for calibration of equipment and Equipment file items such as maintenance;
			Policy and procedure for traceability and required documentation, Policy and procedure for internal audits; Policy and Procedure for data analysis and control of nonconforming
			work; and
	1,3,6,9	2.2.2	Policy for corrective actions (CAs) and preventative actions (PAs).  The organizational chart clearly depicts laboratory structure with
			quality and technical personnel listed.

1, 9  2.2.3 The policy for human resources provisions includes hirin assignment of staff, competence and responsibilities for pand a procedure of training for each technical competence.	
and a procedure of training for each technical competence	ositions,
	e, including
proficiencies required.	uti a a l
1, 3, 4, 2.2.4 Policies for work environment and safety protocols, analy methods, and quality control performed for the National States	
Sanitation Program (NSSP) are included or referenced in	uie QA
Manual and shall be provided upon request.	lab anatam:
1, 9 2.2.5 The policy for sample rejection criteria includes what the will accept and reject based on NSSP requirements and control of the co	
custody.	nani Oi
1, 3, 4, 2.2.6 The laboratory shall have sample acceptance procedures	that include
6, 9 safe handling, transport, and storage to prevent contamina	
deterioration and to protect the sample integrity. These pi	
are provided to customers.	loccuures
1, 3, 4, 2.2.7 The laboratory has procedures for handling nonconforming	na camplac
1, 9 2.2.8 A policy regarding appropriate equipment file maintenan retention (e.g., calibration records, maintenance document	
	itation,
manuals of operation) is included in the QA Manual.  1, 9 2.2.9 The SOP for calibration and maintenance of equipment is	kent or
referenced in the QA Manual and shall be provided upon	
1, 9 2.2.10 The SOP for traceability of analytical results is included on	•
referenced in the QA Manual and shall be provided upon	
This traceability procedure includes a documented procedure	
unique identification of samples and the process for chair	
verification.	1 of custody
1, 9 2.2.11 The QA Manual has a policy and a procedure for internal	l quality
audits. These audits are planned and scheduled annually of	
needed. The policy states auditors do not audit their own	
case of a single person laboratory, FDA LEOs will assist	
audit plan.	
1, 9 2.2.12 The QA Manual contains a policy for data analysis to req	uire that all
analyses performed have been carried out correctly, docu	
controls were used accurately and the results meet specific	
requirements.	
1, 9 2.2.13 The QA Manual contains a procedure for the control of	
nonconforming work in the case of:	
identification, documentation, evaluation, segregation (w	here
practical), disposition of nonconforming sample/analyte/i	
customer notification;	
assigning responsibility for the review and the authority f	for
disposition of nonconforming sample/analyte/result;	
a nonconforming result correction and the re-verification.	/calibration
of the affected equipment after the correction to demonstr	
conformity (if necessary); and	
□ handling a nonconforming result when it is detected, after	r delivery to
the customer.	
1, 9 2.2.14 The QA manual contains a procedure for preventative act	tions in
which laboratory staff identify potential nonconformities	
results, quality records, or customer complaints through a	
process. Steps are then determined to identify preventive	
implement. The necessary changes are made to SOPs and	l this
exercise is recorded, and records maintained.	
1, 3, 6, 9 2.2.15 The QA manual has a policy and a procedure for develop	oing

1			
			corrective action(s) to eliminate the cause of identified
			nonconformities in order to prevent recurrence. Corrective actions
			describe the nonconformities, define the process for evaluating the
			need for actions to ensure that nonconformities do not recur (root
			cause analysis), explain the process to implement the corrective
			action(s) needed, and the resultant outcome. There is also a
			procedure to monitor progress of any ongoing corrective actions and
			the resolution.
	1, 3, 4,	2.2.16	The QA Manual contains a policy stating laboratory management
	6, 9		shall ensure and document the competence of staff independently
			operating equipment resulting in a documented measurement,
			analysis result, quality control value/result, determination of data
			value for sample result, and review/closure of corrective action for
DADELIT C			efficacy.
PAKT III- Qua	nity Contro		nentation for Quality System Defensibility
	1.0		cumentation
	1, 9	3.1.1	The laboratory investigates proficiency testing (PT) programs for
			areas of continual improvement and actively addresses problematic
	1 0 10	212	results through the prescribed corrective action process.
	1, 9, 10	3.1.2	The laboratory personnel performing sampling and testing
			participate in PT programs and exercises when available. If no PT
	126	212	exists, participation in interlaboratory comparisons is considered.
	1, 3, 6,	3.1.3	Corrections to quality control records, bench sheets and reports
	9, 10		follow the requirements below:
			A single line is drawn through the incorrect information;
			The correct information is written next to the incorrect information;
			The person responsible for the correction initialed the information;
			If not obvious, the reason for correction has been included; and
			If corrections are necessary in an electronic document, old
			information must be retained in some form, the person making the
			change must be identified, the date of the change noted, and the
	1 2 6	3.1.4	reason for the change noted.  All records, required to be retained for two years (or length of time
	1, 3, 6, 9, 10	ال.1.4	All records, required to be retained for two years (or length of time as dictated by State law), shall be legible and shall be stored in such
	9, 10		a way that they are readily retrievable to prevent damage or loss.
	1	3.1.5	All records and documents must be written in indelible ink.
	1		thod Performance Validation
	1, 3, 6, 9	3.2.1	The laboratory will internally validate new methods to confirm with
	2, 5, 0, 9	٠.2.1	objective evidence that the intended protocols are demonstrated and
			outcomes are fulfilled.
	1, 9	3.2.2	Methodologies do not deviate from the validated method and the
	-, /		laboratory's internal validation shall remain on file in the laboratory.
	1, 3, 6,	3.2.3	The laboratory shall report the method chosen in writing to the
	9, 10		customer.
	1, 4, 9	3.2.4	Methodologies and protocols are selected based on NSSP
			requirements and samples are processed as per the citation in the
			current Model Ordinance.
	1, 4, 9	3.2.5	Methodologies and protocols are selected based on NSSP
			requirements, and samples are processed as per the citation in the
		<u> </u>	current Guide for the Control of Molluscan Shellfish.
			rironmental Conditions
	1, 3, 4,	3.3.1	Laboratory facilities for analysis, including lighting and
	5, 6, 9,		environmental conditions such as temperature and humidity, shall
	10 1, 3, 4,		support accurate performance of the tests.
l		3.3.2	The laboratory shall monitor, control, and record environmental

5, 6, 9,		conditions as required by the relevant specifications, methods and
10		procedures, or where they influence the outcome of results (e.g.,
		biological sterility, dust, humidity, electrical supply, temperature,
		vibration).
1, 3, 4,	3.3.3	Laboratory personnel shall stop testing when the environmental
6, 9, 10		conditions jeopardize the results of analyses.
1, 3, 4,	3.3.4	Personnel shall ensure good housekeeping in the laboratory.
6, 9, 10		
	3.4 Eq	uipment
1, 3, 4,	3.4.1	The laboratory shall have instructions and/ or SOPs on the use and
6, 9, 10		operation of all relevant equipment, and on the handling and
		preparation of items for testing, where the absence of such could
		jeopardize the outcome of analysis or influence results.
1, 9, 10	3.4.2	All equipment in the laboratory is labelled with the manufacturer's
		name, identification number, and serial number or other unique
		identification that is traceable.
1, 9, 10	3.4.3	Equipment files contain reports and certificates of all calibrations,
		the due date of next calibration, dates and results of any
		maintenance, adjustments, damage, malfunction, and modification or
		repair to the equipment.
1, 9, 10	3.4.4	If equipment (e.g., thermometer, balance) was sent out of the
		laboratory for service, performance has been verified prior to use
		again in the laboratory.
		mperature Measuring Devices
1, 8, 9,	3.5.1	Serial number, ice point date (if applicable) and any correction factor
10		is recorded on in use temperature measuring device (TMD).
1, 8, 9,	3.5.2	TMDs are calibrated as per the NSSP requirements and ice
10		points/steam points are performed annually on Standards
		thermometers.
1, 8,	3.5.3	TMDs calibration certificates are retained for three consecutive
		calibration cycles.
1, 8, 9,	3.5.4	Where calibrations give rise to a set of correction factors, the
10		laboratory shall have procedures to ensure these records are retained
		until the next check is performed.
1, 8, 9,	3.5.5	Range and graduations of all TMDs are appropriate for the
10		designated use. Dial thermometers are not used in the laboratory.
1, 8, 9,	3.5.6	Temperature Monitoring Systems (wired/wireless) must record
10		temperature reading from each sensor/probe in the piece of
		equipment being monitored at the same or greater frequency and
		accuracy as stipulated for mercury in glass thermometers, as per
	2 ( D'	manufacturer specifications.
1 2 4		sposables and Pipettors
1, 3, 4,	3.6.1	Pipettors, accuracy checked, fixed volume or electronic are
6, 9, 10	2.62	calibrated according to NSSP requirements.
1, 3, 10	3.6.2	Pipettors are etched with identification (imprinted serial numbers acceptable) and tagged with last date of accuracy check.
1, 2, 3,	3.6.3	Appropriate pipettor tips are used and sterility checks are performed
4, 6, 9, 10		on an appropriate quantity.
1, 2, 3,	3.6.4	Sterility checks on disposables are performed according to a cited
4, 6, 9,	3.0.4	QC practice, within a designated SOP. (e.g., laboratory may cite and
10		implement a recognized standard of sterility testing, they may test
		10% of a "lot" or any 3 in a box.)
	3.7 Te	st Record/Bench Sheet Requirements
1, 3, 4,	3.7.1	Test records/bench sheets shall contain information to facilitate
 1, 2, 7,	3.7.1	105t 1000tds/00ffcff shoots shall contain information to facilitate

6, 9, 10		repeatability under conditions as close as possible to the original including QC information (or reference) for media and supplies used.
1, 9, 10	3.7.2	Test records/bench sheets must show date, time and temperature of samples at the start of analysis and contain the name or initials of the analyst performing the test for each group of samples.
1, 4, 9, 10	3.7.3	Test records/bench sheets must include sterility controls or a reference to the document containing sterility controls for disposables and dilution buffer.
1, 4, 9, 10	3.7.4	Test records/bench sheets must include media productivity (positive and negative) controls or a reference to the document containing media productivity controls.

## **REFERENCES**

- 1. Good Laboratory Practice.
- 2. U.S. Department of Commerce. 1976. *NBS Monograph 150*. U.S. Department of Commerce, Washington, D.C.
- 3. Association of Official Analytical Chemists (AOAC). 1991. Quality Assurance Principles for Analytical Laboratories. AOAC, Arlington, VA.
- 4. Interstate Shellfish Sanitation Conference (ISSC). 2017. ISSC, Columbia, SC.
- 5. Title 21, Code of Federal Regulations, Part 58, Good Laboratory Practice for Nonclinical Laboratory Study. U.S. Government Printing, Washington, D.C. Technical Programs Criteria for Laboratories Performing Food Testing. AOAC, Arlington, Va.
- 6. U.S. Environmental Protection Agency (EPA). 1975. *Handbook for Evaluating Water Bacteriological Laboratories*. EPA 670/9-75-006. U.S. EPA, Cincinnati, Ohio.
- 7. U.S. Food and Drug Administration (FDA). 1995. *Bacteriological Analytical Manual*. U.S. FDA, 8<sup>th</sup> Edition, AOAC, Arlington, VA.
- 8. National Institute of Standards and Technology Special Publication 250-23, 128 pages (Sept. 1988)U.S. Government Printing office, Washington, D.C. Library of Congress Catalog Number: 88-6000580.
- 9. The International Organization for Standardization and the International Electrotechnical Commission. Online: https://www.iso.org/obp/ui/#iso:std:iso-iec:17025:ed-2:v1:en accessed June 6, 2017.
- 10. National Conference on Interstate Milk Shipments. Cultural Procedures, 2400 Form. Online: http://ncims.org/programs/ accessed June 6, 2017.

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of\_

LAB	ORATO	ORY:	DATE of EVALUATION:				
	SHELLFISH LABORATORY EVALUATION CHECKLIST						
		SUMMARY of NONCONFORMITI	ES				
Page	Item	Observation	Documentation Required				
	<u> </u>						

Revised 6-16-2017

LABOR.	ATORY STATUS	
LABOR.	ATORY	DATE
LABOR	ATORY REPRESENTATIVE/POINT OF CONTACT:	
	NSSP Quality System Evalua	tion: (Part I-III)
A. Cı	riteria for Determining Laboratory Status of the Qualit	y System Component:
1.	Laboratory must satisfy all sections of the Quality S	ystem prior to onsite evaluation:
	a. The total # of nonconformities in Part I	
	b. The total # of nonconformities in Part II	
	c. The total # of nonconformities in Part III	
B. La	aboratory Status (circle appropriate)	
Do	oes Not Conform	Conforms
Acknowl	ledgment by Laboratory Director/Supervisor:	
All Corre	ective Actions will be implemented and verifying substantia	ating documentation received by the Laboratory
Evaluatio	on Officer on or before	so onsite evaluation can be scheduled.
Laborato	ory Signature:	Date:
LEO Sigi	nature:	Date:

LAB	LABORATORY:					
Page	Item	Observation				