Hazard Analysis and Critical Control Point
Training Curriculum

Developed by the National Seafood HACCP Alliance
for Training and Education
National Seafood HACCP Alliance

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Seafood HACCP Training

The National Seafood HACCP Alliance and the Association of Food and Drug Officials (AFDO) have developed courses for training in basic HACCP programs and related Sanitation Control Procedures. Train-the-Trainer courses are also available.

For the most current course information, please consult:
Seafood Network Information Center (http://seafood.ucdavis.edu)
Association of Food and Drug Officials (http://afdo.org)

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National Seafood HACCP Alliance for Training and Education

The National Seafood HACCP Alliance (SHA) provides a current, convenient and cost-effective education and training program to assist commercial and regulatory compliance with the prevailing requirements for product safety during processing and import of any seafood in the United States. The structure for the SHA program is based on collaboration among federal and state food-inspection officials, academic food safety researchers and educators, and various representatives from the seafood and aquaculture industry with commerce in the U.S. (Slide 1). The program is directed by a voluntary SHA Steering Committee. Any individual, company, agency or nation can participate in the SHA program through communications with the SHA Steering Committee. The SHA Steering Committee directs development of all training materials and courses, and the accompanying SHA Protocol.

The SHA Protocol is maintained by the Association of Food and Drug Officials (http://www.AFDO.org). SHA Protocol are intended to maintain a uniform and standard training format based on qualified trainers, current training materials, approved courses, and course audits for both domestic and international audiences (Slide 2). AFDO records and issues all certificates for participants that complete an approved SHA course. Certificates are issued for SHA courses in 'HACCP: Hazard Analysis and Critical Control Point Training' and 'Sanitation Control Procedures for Processing Fish and Fishery Products.' The HACCP course is offered in both a formal classroom setting and through a self-guided Internet version that participants can complete online. To obtain an AFDO certificate of course completion, the Internet participants must later attend a one-day “Segment Two” session in person.

The SHA training materials include the standard training manuals, model HACCP plans, the FDA Hazards Guide and Compendium of Fish and Fisheries Product, Processes, Hazards, and Controls. The HACCP models are for reference as example plans for processing and importation of various seafood products (Slide 3). Some of these materials have been translated for international audiences.
The SHA program and training materials are designed to meet the HACCP training requirements established under Title 21 Code of Federal Regulations (CFR) Part 123.10 of the U.S. Food and Drug Administration’s mandatory seafood HACCP inspection program. The mandates in 21 CFR Part 123.10 require that certain HACCP activities must be completed by a “HACCP-trained individual.” The National Seafood HACCP Alliance course is the standardized curriculum by which FDA will evaluate other training courses (Slide 4). A HACCP-trained individual is one who has successfully completed FDA-recognized training in the application of HACCP to fishery products (at least equivalent to that received under a “standardized curriculum” recognized by FDA) or has acquired the knowledge through job experience. Attending an SHA course is not mandatory, but it does provide assurances that the resulting knowledge will be consistent with regulatory expectations.
History

The SHA program began as an idea during the April 1993 National Sea Grant Forum on Seafood Safety and Quality. In conjunction with the Association of Food and Drug Officials of the Southern States (AFDOSS), their Board of Directors passed a resolution to advance a seafood HACCP training program. The Council of Sea Grant Directors followed with financial support for the first meeting of the National Seafood HACCP Alliance held in December 1993. Since this modest beginning, the SHA program has received continuing support through grants from FDA, the U.S. Department of Agriculture (USDA) National Institute of Food and Agriculture (formerly the Cooperative State Research, Education, and Extension Program) and the National Sea Grant College Program (Slide 5). Although there are many HACCP training programs and consultants, the SHA training program remains distinct in that it is the primary and proven training program recognized by all pertinent seafood regulatory authorities in the United States. It is the foundation training program for most regulatory agencies monitoring seafood commerce in the United States.

Program support for the Seafood HACCP Alliance has been provided through grants from:
• National Sea Grant College Program
• United States Food and Drug Administration (FDA)
• United States Department of Agriculture – National Institute of Food and Agriculture (USDA NIFA)

The SHA program is now sustained, in part, by the program fees and the generous, voluntary labor of the SHA Steering Committee and qualified trainers about the world. Through 2009, nearly 25,000 participants have completed SHA HACCP courses conducted in every seafood producing nation in the world. Participation has included individuals from both government and commercial operations addressing all aspects of seafood and aquaculture products from production and processing through distribution, wholesale, retail and food services/restaurants. The SHA approach recognizes the essential role of regulatory authorities, the educational networks of Sea Grant and the Cooperative Extension Service, and the need for regional and international programs.

Maintaining Course Integrity

Because this course will be used to evaluate HACCP-training equivalency, it is imperative that course instructors adhere to the course format and material to the extent possible. The course is divided into three segments. The first teaches the student the seven principles of HACCP. The second segment explains the seafood HACCP regulations and guidance materials available to help develop a HACCP plan. The last segment is a class exercise where students are divided into small groups and asked to conduct a hazard analysis and develop a HACCP plan. Each of these segments is necessary to give students an adequate foundation to establish their firm’s HACCP mandate.
HACCP Course Agenda

The course agenda is a suggested outline of how the course may be structured. The standard agenda is a 2.5 day (20 hours) course. The times allotted to each section are recommendations to allow for sufficient learning opportunities. However, there is flexibility in the design based on the nature of the audience (i.e., same processing, homogeneous audience, smaller size). Regardless of the format of the course, it is important that all agenda items be covered and that 3-4 hours be allowed for the practical work session. One approach to stimulate participation is to arrange the work sessions following the respective instruction (e.g., the work session on hazard analysis to follow the lecture on Determining Critical Control Points, and the work session on Developing the HACCP Plan following the lecture on Recordkeeping.) Likewise, some may choose to teach the HACCP regulation on day one, prior to HACCP principles. Instructors may also elect to supplement information in Chapter 4 (Seafood Safety Hazards) with additional seafood-specific hazards materials unique to the audience, product types or region.

Day One

20 minutes Welcome and Course Objectives Chapter 1
60 minutes Prerequisite Programs Chapter 2
20 minutes Preliminary Steps HACCP Chapter 3
75 minutes Seafood Safety Hazards Chapter 4
90 minutes Hazard Analysis Chapter 5

Lunch

60 minutes Determine Critical Control Points Chapter 6
60 minutes Establish Critical Limits Chapter 7
40 minutes Critical Control Point Monitoring Chapter 8
40 minutes Corrective Actions Chapter 9

Day Two

60 minutes Establish Verification Procedures Chapter 10
60 minutes Record-Keeping Procedures Chapter 11
90 minutes The Seafood HACCP Regulation Chapter 12

Lunch

30 minutes Sources of Information on Preparing HACCP Plans Chapter 13
60 minutes Review and Preparation for Developing HACCP Plans
<table>
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<td>Work Sessions on Conducting Hazard Analysis (break into groups)</td>
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<tr>
<td>90 min</td>
<td>Work Sessions on Developing HACCP Plans (break into groups)</td>
</tr>
<tr>
<td>30 min</td>
<td>Present Results from Work Sessions</td>
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Introduction to Alliance Course and HACCP

Course Objective

This course provides training for the seafood industry and regulators consistent with regulatory requirements of the December 1995 United States Food and Drug Administration (FDA) Seafood Hazard Analysis and Critical Control Point (HACCP) regulation (see Chapter 12 and Appendix 1). The regulation became effective in December 1997 to ensure safe processing and importing of seafood products in the United States. This regulation specifies that training is required for persons responsible for developing and modifying the HACCP plan and reviewing records. This course contains the information necessary for you or other members of your team to meet the HACCP training requirements outlined in the FDA's regulation (Slide 1). Numerous state authorities and nations have adopted various versions of HACCP for seafood safety.

Slide 1

In this chapter, you will learn the:

• Objective of the course.
• Format of the course.
• Expectations of the participants, and
• Meaning and importance of HACCP.

Course Format

This seafood HACCP course is divided into three distinct parts:

• HACCP fundamentals,
• Explanation of HACCP and FDA's regulation and guidance materials to help develop a HACCP plan, and
• Class exercises to provide practice and instruction in the development of seafood HACCP plans.
The first part defines the seven principles of HACCP. Learning these principles will give a better understanding of the fundamentals on which HACCP is based. As each principle is discussed, the class will progressively develop a HACCP plan for a common fresh fish, mahi-mahi, produced in fillet form by the fictional model “XYZ Seafood Company”. This example will help you understand HACCP principles and how they relate to seafood processing.

The example incorporates the use of the FDA *Fish and Fisheries Products Hazards and Controls Guidance* and provides information about seafood-specific hazards.

The second part explains the seafood HACCP regulations and guidance materials that are available to help you develop a HACCP plan. This training manual also provides information about seafood-specific hazards.

The third part is a practical exercise that demonstrates how to develop a seafood HACCP plan. During this part of the course, the class will be divided into teams to write HACCP plans for different seafood products.

**What is Expected of the Participant**

HACCP is an important safety-management system and can be integrated into any operation. However, HACCP can seem complicated and demanding until the basic concepts are understood. Therefore, you are encouraged to ask questions and to contribute first-hand experiences during the discussions. This manual includes exercises that require class participation throughout the training. Keep in mind that the more you contribute to these exercises, the less complicated the HACCP system will seem and the easier it will be to develop and implement a HACCP plan.

**How to Use This Training Manual**

This manual is yours. Become familiar with it. Learn where the definitions are, where the forms are that will help you develop a HACCP plan, and where to find other basic information. Make as many notes and marks in the text as needed to assist in creating and understanding a HACCP plan. Use the manual as a reference. This manual does not have a copyright. Make as many copies of the enclosed forms as necessary or copy the whole manual to share with others in your company.

**Meaning and Importance of HACCP**

HACCP is an acronym that stands for Hazard Analysis and Critical Control Point (Slide 2). It is a systematic approach to the identification, evaluation and control of food-safety hazards. The concepts behind this term are important.

---

**Slide 2**

HACCP stands for **Hazard Analysis and Critical Control Points**.
HACCP is a preventive system of food safety hazard controls rather than a reactive one (Slide 3). Food processors can use it as a management tool to ensure safer food products for consumers. The HACCP system is designed to identify hazards (HA – Hazard Analysis) and establish controls (CCP – Critical Control Points).

HACCP concepts are:
• Preventive, not reactive.
• A management tool used to protect the food supply.

HACCP is not a zero-risk system, but it is designed to minimize the risk of food-safety hazards to acceptable levels (Slide 4). It is a proven approach to help assure food safety. In 1973, FDA required HACCP-type controls for processing low-acid canned foods to protect against *Clostridium botulinum*, the bacteria that can produce the toxin which causes botulism.

The HACCP concept was first applied to food production during efforts to supply safe food for the United States space program in the early 1960s. It was decided that existing quality control techniques did not provide adequate assurance against contamination during food production. The end-product testing necessary to provide assurance that the food would be safe would be so extensive that little food would be available for space flights (Slide 5). Currently, HACCP has been adopted by many food processors to assure the safety of the food products.

Origins of HACCP:
• Pioneered in the 1960s.
• First used when foods were developed for the space program.
• Adopted by many food processors.
In 1985 the National Academy of Sciences (NAS) recommended that the HACCP approach be adopted by all regulatory agencies and that it should be mandatory for food processors. This recommendation led to the formation of the National Advisory Committee on Microbiological Criteria for Foods (NACMCF). This committee standardized the HACCP principles used by industry and regulatory authorities. The committee’s work is the basis of this core curriculum (Slide 6).

**Slide 6**

National Academy of Sciences recommendation:
The HACCP approach should be adopted by all regulatory agencies and it should be mandatory for food processors.

HACCP is endorsed worldwide by many countries and organizations such as Codex Alimentarius (a commission of the United Nations). Although the regulatory approaches may differ from one nation to another, the HACCP concepts are the same (Slide 7).

**Slide 7**

Examples of international use:
- Codex Alimentarius
- Australia
- Canada
- Chile
- China
- Ecuador
- European Union
- Japan
- New Zealand
- South Africa
- many other nations

FDA’s Seafood HACCP regulation and other domestic and international HACCP control systems are based on seven basic principles or steps (Slide 8). The principles will be explained in more detail in this course.

**Slide 8**

Seven principles of HACCP:
1) Conduct a hazard analysis.
2) Determine the critical control points (CCPs) in the process.
3) Establish the critical limits.
4) Establish monitoring procedures.
5) Establish corrective actions.
6) Establish verification procedures.
7) Establish record-keeping procedures.
HACCP is a preventive system for ensuring food safety, but it is not a stand-alone system. To be effective, HACCP must be built upon current food safety programs such as Good Manufacturing Practices (GMPs) and Sanitation Standard Operating Procedures (SSOPs). These programs are known as “prerequisites” that provide a foundation for the HACCP program (Slide 9).

As you learn more about HACCP, there will be many new definitions that you will need to understand. To assist you, the most common HACCP definitions are found in the following two pages. Refer back to these pages as needed and add other terms as appropriate that will help you in developing and implementing your own HACCP plan.

**Definitions**

**CCP:** Critical Control Point

**CCP Decision Tree:** A sequence of questions asked to determine whether a control point is a CCP.

**Continuous Monitoring:** Uninterrupted collection and recording of data such as time-temperature recorder.

**Control:** (a) *verb* To manage the conditions of an operation in order to maintain compliance with established criteria. (b) *noun* The state in which correct procedures are being followed and criteria are being met.

**Control Measure:** Any action or activity that can be used to prevent, eliminate or reduce a significant hazard (previously known as a preventive measure).

**Control Point:** Any point, step or procedure at which biological, physical or chemical factors can be controlled.

**Corrective Action:** Procedures followed when a deviation occurs.

**Critical Control Point (CCP):** A step at which control can be applied and is essential to prevent or eliminate a food-safety hazard or reduce it to an acceptable level.

*Source of Definitions*

**Critical Limit:** A maximum and/or minimum value to which a biological, chemical or physical parameter must be controlled at a CCP to prevent, eliminate or reduce to an acceptable level the occurrence of a food-safety hazard.

**Deviation:** Failure to meet a critical limit.

**HACCP:** Hazard Analysis and Critical Control Point

**HACCP:** A systematic approach to the identification, evaluation and control of food-safety hazards.

**HACCP Plan:** The written document that is based upon principles of HACCP and that delineates the procedures to be followed.

**HACCP System:** The result of the implementation of the HACCP plan.

**HACCP Team:** The group of people who are responsible for developing, implementing and maintaining the HACCP system.

**Hazard:** A biological, chemical or physical agent that is reasonably likely to cause illness or injury in the absence of its control.

**Hazard Analysis:** The process of collecting and evaluating information on hazards associated with the food under consideration to decide which are significant and must be addressed in the HACCP plan.

**Monitoring:** To conduct a planned sequence of observations or measurements to assess whether a CCP is under control and to produce an accurate record for future use in verification.

**Operating Limits:** Criteria that are more stringent than critical limits and that are used by an operator to reduce the risk of a deviation.

**Prerequisite Programs:** Procedures, including Good Manufacturing Practices (GMPs), that address operational conditions providing the foundation for the HACCP system.

**Severity:** The seriousness of a hazard (if not properly controlled).

**Validation:** The element of verification focused on collecting and evaluating scientific and technical information to determine if the HACCP plan and/or processes employed, when properly implemented, will effectively control the hazards.

**Verification:** Those activities that determine the validity of the HACCP plan and that the system is operating according to the plan.
Prerequisite Programs

Introduction to Prerequisite Programs

HACCP is not a stand-alone program, but part of a larger system of control procedures to ensure food safety. For HACCP to function effectively it needs to be accompanied by what are called “prerequisite programs” (Slide 1).

Slide 1

In this chapter, you will learn:
• The importance of prerequisite programs for HACCP,
• Good Manufacturing Practices (GMPs),
• Sanitation Control Procedures (SCPs), and
• Examples of SCP monitoring frequency.

Prerequisite programs provide the basic environmental and operating conditions that are necessary for the production of safe and wholesome food. Some of these programs are required by regulations and others are recommended (Slide 2).

Slide 2

Definition:
Prerequisite programs are procedures, including Good Manufacturing Practices (GMPs), that address environmental and operational conditions which provide the foundation for the HACCP system.
Good Manufacturing Practices (GMPs)

Good Manufacturing Practices (GMPs) are federal regulations (FDA, Code of Federal Regulations, Title 21, Part 110 – see Appendix 3) that apply to all food manufacturers and are the basis for determining whether the process facility, methods, practices and controls used to process food products are safe and whether the products have been processed under sanitary conditions. GMPs outline the minimum standards that a food processing facility needs to meet including (but not limited to): personnel, buildings and facilities, equipment, production and process controls, raw materials, and manufacturing operations (Slide 3).

**Web Link**
An Internet training course on the basic requirements of the FDA Good Manufacturing Practices regulation is available at http://gmptraining.aem.cornell.edu

**Web Link**
Sanitation Control Procedures for Processing Fish and Fishery Products (SGR 119) is available through the University of Florida IFAS Extension Bookstore, http://www.ifasbooks.com (follow the HACCP link)

Sanitation Control Procedures (SCPs)

HACCP systems must be built upon a firm foundation of compliance with the GMPs and acceptable sanitation control procedures (SCPs). (Slide 4)

SCP are the necessary procedures to meet GMPs requirements. When SCPs are in place, HACCP can be more effective because it can then concentrate on the hazards associated with the food or processing and NOT the processing plant environment or employee practices (Slide 5). SCPs monitoring and corrections shall be included to meet the record requirements in the seafood HACCP regulation. Written SCPs are recommended to outline the goals of the sanitation program and how it will be implemented.

**Slide 3**
Good Manufacturing Practices (GMPs) are the basis for determining if processing methods are safe and food is being processed under sanitary conditions.

**Slide 4**
HACCP program must be based on a solid foundation in compliance with the GMPs and SCPs.

**Slide 5**
Sanitation control procedures (SCPs) are used by food processing firms to meet requirements in the GMPs.

SCP are an effective means to control potential food safety hazards that might be associated with the processing environments and employee practices.
Well designed SCPs that are properly implemented are an effective means to control food safety hazards that might be associated with the processing environment and employee practices. For example, SCPs can be designed to help control some bacterial hazards by specifying procedures to:

- Prevent cross contamination by maintaining proper product flow and limiting certain employee tasks and movement.
- Prevent contamination by employees by locating hand washing and sanitizing stations in the processing area to better facilitate proper hand washing.
- Prevent contamination by using proper cleaning and sanitizing procedures for equipment.

Likewise SCPs can also be used to help control some chemical contamination hazards by specifying procedures to:

- Prevent contamination by storing chemicals in an appropriate place.
- Prevent contamination by labeling all chemicals.
- Prevent contamination by using chemicals per label directions.

By properly implementing these types of SCPs, a processor is able to control potential food safety hazards that may be associated with the plant environment and employee practices outside of a HACCP plan.

**Additional Prerequisite Programs**

In addition to GMPs and SCPs there are other prerequisite programs that a processor may need to have in place in order to support the HACCP program. (Slide 6).

**Slide 6**

Additional prerequisite programs:
- Employee training
- Supplier controls
- Traceability and product recall procedures
- Preventive maintenance

- **Employee training**: Employee training programs are an essential component of an effective food safety and sanitation program. Employees need to know how to perform their job and understand how their work can impact the safety of the product.

- **Supplier controls**: Procedures must be in place to ensure that suppliers have effective GMP, SCP, HACCP, or other food-safety programs in place. Written product and packaging material specifications should be developed for all suppliers. The product specifications should include a description of any potential biological, chemical and physical hazards that might need to be controlled as well as any quality requirements that have been determined.
• **Traceability and product recalls:** Procedures should be in place to ensure that raw material and finished products are coded and identified properly in order to trace the product backward (from receipt) and forward (into distribution). Traceability procedures are required by the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 (Bioterrorism Act). A product recall system should also be in place so that products can be removed from the marketplace quickly when necessary. Recall procedures can be found in Code of Federal Regulation (CFR) 21, Part 7, Enforcement.

• **Preventive maintenance:** Facility and equipment often require routine maintenance before problems develop. Preventive maintenance schedules and procedures should be established and documented using manufacturer and other information as needed.

Finally, there are a number of requirements that are also considered prerequisite programs that a processor should know (Slide 7). This course cannot discuss all prerequisite program in detail. Depending on the product or business, there will be additional compliance requirements that apply to the foods being processed.

**Slide 7**

Other requirements that are considered prerequisite programs:
- Local regulations
- Food defense and biosecurity requirements – Department of Homeland Security (DHS)
- Allergen labeling requirements – Food Allergen Labeling and Consumer Protection Act of 2004
- Country of origin labeling (COOL) requirements – 2002 Farm Bill
- Nutritional labeling – Nutritional Labeling and Education Act, 1994 (NLEA) Requirements
- Regulatory food standards
- Procedures to guard against economic fraud

There may be specific state or local code requirements for food processing establishments in your area that specify where your operation should be located, and how it is constructed and maintained. You may also need to obtain specific permits or licenses from state or local authorities. There may be product-specific requirements for nutritional and allergen labeling. If the product is being imported, it is necessary to meet the requirements in the Bioterrorism Act.

**FDA Eight Key Sanitation Areas**

FDA’s seafood HACCP regulation focuses on eight key areas of sanitation (Slide 8). These eight areas of sanitation are derived from the GMPs (see Appendix 3). FDA’s seafood HACCP regulation does not require that SCPs program be written. However, FDA requires that these eight key areas of sanitation be monitored and that the results be recorded along with any corrections for deficiencies.
Eight key areas of sanitation:
1) Safety of water
2) Condition and cleanliness of food contact surfaces
3) Prevention of cross contamination
4) Maintenance of hand washing, hand sanitizing and toilet facilities
5) Protection from adulterants
6) Labeling, storage and use of toxic compounds
7) Employee health
8) Exclusion of pests

Not all of these eight areas may be relevant to all processing facilities. For example, a processor that receives frozen fish, stores it frozen and ships it frozen may not need to address the condition and cleanliness of food contact surfaces. Monitoring is not required for those key areas that are not relevant. The eight key areas were designed to include those aspects of the GMPs that are most likely to have an impact on the product safety.

A general description of each of the eight key areas of sanitation is provided below.

1) **Safety of water:** Water (and ice) that contacts food or food-contact surfaces shall be of safe and of sanitary quality (Slide 9)

   • Source of water, and the plumbing system that conveys it to the building must provide a safe supply. For companies using a municipal water source for processing and making ice, the water treatment authority is responsible for safety of source and conveyance to the building. Documents must include annual water quality tests from the water authority. Companies using private water systems (e.g. wells), are directly responsible for adequate monitoring and documentation of the safety of the water source. Municipalities can provide guidance.
   • Protection of water used for food and food manufacturing must prevent contamination from potential cross connections and backflow. To ensure water is safe, cross-connections must be prevented. There must be no cross connection or backflow potential between the water supply and piping for wastewater or sewage.
2) **Condition and cleanliness of food contact surfaces:** Food contact surfaces shall be of a proper design and maintained in a clean and sanitary manner to prevent food contamination (Slide 10).

- Food contact surfaces must be designed, fabricated, maintained, and installed so that they are easy to clean and able to withstand the environment in which they are used. Equipment must have smoothly bonded seams and be made of impervious materials that can be easily cleaned and sanitized. Food contact surfaces must also be maintained in suitable condition. For example, poorly bonded joints, corroded parts, exposed bolts, screw heads, or even pitted surfaces that could trap water or soils should not be used.
- Adequate cleaning and sanitizing procedures and frequencies must be established for all food contact surfaces, including equipment, utensils, food containers, gloves, and outer garments. Suggested frequencies for cleaning and sanitizing include: before use, after processing interruptions, and as necessary. Cleaning is accomplished using detergent and water at a suitable temperature. Sanitizing is accomplished using approved sanitizing agents such as chlorine, quaternary ammonium or iodine-based compounds.

3) **Prevention of cross contamination:** Employee hygiene, personnel practices and the design of the facility must prevent cross contamination (Slide 11).

- Employees shall maintain adequate personal cleanliness and conform to adequate hygienic practices to prevent product contamination. Workers must wear clean and appropriate attire and must wash and sanitize their hands at appropriate intervals. Gloves must be used appropriately and are not a substitute for hand washing and sanitizing.
- Employees must understand that their actions can contribute to product contamination. Employees’ hands or gloves, along with equipment and utensils must be washed and sanitized (when necessary) after being contaminated. For example, employees working in the raw product area should not work with the finished product without washing and sanitizing their hands, gloves, equipment, or utensils to avoid cross-contamination.
- Plant design must prevent potential contamination of food, food contact surfaces, and packaging material. Operations must be adequately separated where contamination is likely to occur. Raw product and unpackaged cooked ready-to-eat product must be separated to avoid contamination. Food contact surfaces must be cleaned and sanitized when contaminated. Packaging materials must be stored and handled properly so they do not become a source of contamination.
3) Prevention of cross-contamination:
- Employee hygiene practices
- Employee food handling practices
- Plant design and layout
- Physical separation of raw and ready-to-eat products

4) **Maintenance of hand washing, hand sanitizing and toilet facilities:**
Sanitary facilities must be accessible, properly maintained, and adequately supplied. An adequate sewage disposal system must be in place (Slide 12).

- Hand washing and, where appropriate, hand sanitizing facilities should be at each location where good sanitary practice requires their use. Effective hand washing and sanitizing supplies must be available. Water at suitable temperature and sanitary towel service or suitable drying devices must be available and designed to prevent recontamination.
- An adequate sewage disposal system is required. Adequate and readily accessible toilet facilities with self-closing doors to protect food from airborne contamination must be maintained in sanitary condition. All toilet facilities must be adequate, in good repair (e.g. not leaking), and properly supplied with paper towels, soap, etc.

5) **Protection from adulterants:** Food, food contact surfaces, and food packaging material must be protected from microbiological, chemical and physical contaminants (Slide 13).

- Potential contamination sources could include: water splashing from the floor, condensate from air conditioners, refrigerator condensers, pipes, light fixtures and ceilings, toxic substances (e.g., pesticides, fuel, cleaning compounds, and sanitizing agents), filth, and physical contaminants (e.g., glass, metal fragments, dirt or corrosion from fans and other fixtures).
6) **Labeling, storage and use of toxic compounds**: Prevent contamination from toxic compounds. Toxic cleaning compounds, sanitizing agents and pesticides must be properly labeled, used and stored in a manner that protects food, food contact surfaces and packaging material from contamination (Slide 14). Toxic compounds must be stored in a secured area with limited access separated from food processing and areas where food and packaging materials are stored.

7) **Employee health**: Food handlers with an apparent illness, wound, or open lesions could be a source of microbiological contamination (Slide 15). Policies must be in place that exclude or restrict employees who exhibit or are diagnosed with symptoms of an illness, wounds or other afflictions that could be a source of microbial contamination of food, food contact surfaces, and food packaging material.

8) **Exclusion of pests**: Pests, such as rodents, birds, domestic animals and insects are not allowed in any area of a food processing and/or storage facility (Slide 16). Even if pest control is contracted to an outside company, the processor is responsible to assure there are no pests in the facility.

**Sanitation requirements**

SCPs are required for the eight key sanitation areas in the Seafood HACCP regulation that apply to each processor’s situation (Slide 17). Written Sanitation Standard Operating Procedures (SSOP) are not a regulatory requirement, but they are recommended. An example of a written SSOP is provided in Table 1 (page 19). Successful implementation of SCPs depends on having adequate procedures in place, effective monitoring and proper corrections. Monitoring results for the eight key areas of sanitation must be recorded. In
addition, when monitoring reveals a deficiency, corrections must be taken and recorded. Table 2 (page 26) shows the relation between the Seafood HACCP Regulation (21 CFR 123.11 (b)) and the current GMP Regulation (21 CFR 110).

Examples of monitoring frequencies and corrections are illustrated in Slide 18. Frequency of monitoring and the necessary corrections depend on the unique situations in each facility. In certain processing plants, the safety of the processing water may be checked annually. However, plants using private water sources such as well water will likely require more frequent monitoring checks.

**SCP Records**

The SCP record information of the eight key sanitation areas is expected by inspectors and should be included in an effective sanitation monitoring program. Required elements of SCP monitoring are listed in Slide 19. The monitoring forms or records are subject to record-keeping requirements outlined in the FDA's seafood HACCP regulation and must include the name and location of the processor, the date and time the monitoring was performed, and corrections made. Each of the eight key areas applicable to the operation must be included. The frequency or time for monitoring will vary according to various types of products and the schedule of operations. Examples of typical SCP monitoring forms are provided in Forms 1-3 (see pages 27-30). Form 1 includes a daily monitoring frequency for both raw and ready-to-eat food. Form 2 illustrates less frequent or periodic monitoring for conditions or situations that are not expected to be frequent problems. Form 3 is an example for a process operation such as a warehouse that has only certain key sanitary concerns.

**Sanitation Controls and HACCP**

Sanitation controls are not typically included in the HACCP plan. Sanitation controls address the overall processing plant environment and employee practices. HACCP controls address specific hazards in the products and processing steps.

For example, hand washing is a general employee practice and not specifically related to the product or processing step. Likewise, routine cleaning of food contact surfaces is not specifically related to a single product or a particular processing step. Examples of hazards that are controlled by a HACCP plan or Sanitation Control Procedure are shown in Slide 20.
Examples of monitoring frequency and corrections

<table>
<thead>
<tr>
<th>Sanitation Condition/Practice</th>
<th>Frequency of Monitoring</th>
<th>Corrections</th>
</tr>
</thead>
</table>
| Safety of water               | Municipal source: Annually  
Private well: Semi-annually  
Cross connections: Semi-annually (unless changes are made) for hard plumbing between potable and non-potable lines  
Cross connections: daily, if hose bibs not protected | Example: If report of water shows high coliform counts, stop processing. Resample water and/or ice to determine required corrections before restarting. |
| Condition and cleanliness of food contact surfaces | Condition of processing equipment: Monthly or more often if equipment is repaired or replaced to assure it meets the construction standards.  
Cleaning and sanitizing of equipment, utensils, gloves, and outer garments that come in contact with food: Daily, every time the equipment is cleaned and sanitized. Raw seafood, once a day at start. Ready-To-Eat (RTE) sea foods, start and every 4 hours  
Record sanitizer concentrations. | Example: If sanitizer concentration is too low, stop. Make new sanitizing agent and clean and sanitize again. |
| Prevention of cross contamination | Plant design: Monthly or more often if modifications are made to the facility.  
Employee practices: Daily, at start of production and at least every four hours during production. More often if necessary to ensure that employees hands, gloves, equipment and utensils are washed and sanitized (as necessary) after being contaminated.  
Separation of raw and cooked products performed daily.  
Coolers and processing area every four hours during operations and at the end of processing to ensure that unpackaged cooked product is separated from raw product. | Example: If raw product touches or otherwise contaminates cooked product, the cooked product will not be distributed and source of problem will be corrected. |
<table>
<thead>
<tr>
<th>Sanitation Condition/Practice</th>
<th>Frequency of Monitoring</th>
<th>Corrections</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintenance of hand washing, hand sanitizing and toilet facilities</td>
<td>Hand washing and hand sanitizing facilities: Daily, during pre-op inspection to make sure soap, warm water and paper towels are available and toilet facilities are in good repair. The concentration of hand sanitizing solutions should be monitored at pre-op and every four hours during processing for RTE products.</td>
<td>Example: If toilet seal is leaking, it must be fixed.</td>
</tr>
<tr>
<td>Protection from adulterants</td>
<td>Protection from adulterants: Daily at start-up and every four hours to make sure food is protected from contaminants like condensate, floor splash, and glass.</td>
<td>Example: Condensation falling from the ceiling, pipes and cooling coils above food handling areas can drip onto food, packaging materials or food contact surfaces. Affected product must be segregated and evaluated, fix insulation of pipes and increase air circulation in the room.</td>
</tr>
<tr>
<td>Labeling, storage and proper use of toxic compounds</td>
<td>Labeling, storage and proper use of toxic compounds: Daily, during pre-op inspection to make sure toxic compounds are properly labeled and stored.</td>
<td>Example: If a bottle is unlabeled, remove the bottle from processing area, identify its content and label appropriately or destroy.</td>
</tr>
<tr>
<td>Employee health conditions</td>
<td>Employee health conditions (illness, wounds, etc.): Daily, before production starts.</td>
<td>Example: Employee who has an infected wound in the hand could be assigned to an area away from food processing.</td>
</tr>
<tr>
<td>Exclusion of pests</td>
<td>Exclusion of pests: Monthly for outside monitoring. Daily monitoring, during pre-op inspection for processing and storage areas. Grounds around a plant may require monthly checks to discourage attraction of pests: checking the inside of the processing facility for pest activity would be daily.</td>
<td>If rodent excrement found, remove and clean area before start. If daily problem, call pest control company and look for source of entry.</td>
</tr>
</tbody>
</table>
Chapter 2

Slide 19

Required Elements of SCP Monitoring Records
• Name and address of the firm
• Date and time of the recorded activity
• Include all of the eight key sanitary concerns pertinent to the operation
• Monitoring procedure and appropriate frequency
• Monitoring results
• Corrections taken
• Signature or initials of person conducting the monitoring

Slide 20

<table>
<thead>
<tr>
<th>Hazard</th>
<th>Control</th>
<th>Type of Control</th>
<th>Control Program</th>
</tr>
</thead>
<tbody>
<tr>
<td>Histamine</td>
<td>Time and temperature controls for fish</td>
<td>Product specific</td>
<td>HACCP</td>
</tr>
<tr>
<td>Pathogen survival</td>
<td>Time and temperature controls for smoking fish</td>
<td>Processing step</td>
<td>HACCP</td>
</tr>
<tr>
<td>Contamination with pathogens</td>
<td>Wash hands before touching product</td>
<td>Employee</td>
<td>Sanitation or SCP</td>
</tr>
<tr>
<td>Contamination with pathogens</td>
<td>Limit employee movement between raw and cooked areas</td>
<td>Employee</td>
<td>Sanitation or SCP</td>
</tr>
<tr>
<td>Contamination with pathogens</td>
<td>Clean and sanitize food contact surfaces</td>
<td>Plant environment</td>
<td>Sanitation or SCP</td>
</tr>
<tr>
<td>Chemical contamination</td>
<td>Use only food-grade grease</td>
<td>Plant environment</td>
<td>Sanitation or SCP</td>
</tr>
</tbody>
</table>
Example of written sanitation standard operating procedures (SSOP)

It is recommended that companies create and implement Sanitation Control Procedures to enable them to meet the documentation requirements. A Sanitation Control Procedures or SCP is a set of procedures a firm will follow that addresses the environmental conditions of the facility.

The following is an example of a written SSOP for a fictitious company producing raw and cooked RTE seafood products:

**Table 1. Model Sanitation Standard Operating Procedure**

1) **Safety of water (FDA Key Sanitation Condition No. 1)**

Controls and Monitoring:

a) All water used in the plant is from a reliable municipal water system. Municipal water bills indicate that the water source is safe. **Monitoring Frequency: Annually.**

b) The water system in the plant was designed and installed by a licensed plumbing contractor, and meets current community building codes. All modifications to the plumbing system will be completed by a licensed plumbing contractor and will be inspected to ensure conformance with local building codes. Copies of building inspection reports indicate that the plumbing system is properly constructed. **Monitoring Frequency: When plumbing is installed or modified.**

c) All water faucets and fixtures inside and outside the plant have antisiphoning controls. Water faucets and fixtures are inspected for the presence of antisiphoning controls. **Monitoring Frequency: Daily before processing.**

Corrections:

a) In the event of municipal water treatment failure, the plant will stop production, determine when the failure occurred, and hold products produced during the failure until product safety can be assured. Production will resume only when water meets state and federal water quality standards.

b) Corrections will be made to the plumbing system, if necessary, to correct problems. Production will resume only when water meets state and federal water quality standards.

c) Water faucets and fixtures without antisiphoning controls will not be used until antisiphoning controls have been implemented.

Records:

a) Municipal water bill and monthly sanitation control record.

b) Building plumbing inspection report and periodic sanitation record.

c) Daily Sanitation Control Record.
Chapter 2

2) Condition and cleanliness of food contact surfaces (FDA Key Sanitation Condition No. 2)

Controls and Monitoring:

a) Food contact surfaces are readily cleanable (do not have cracks, cavities, crevices, overlapping joints, mineral scale, etc. that are not possible to adequately clean and sanitize). The sanitation supervisor inspects food contact surfaces to determine if they are readily cleanable. **Monitoring Frequency: Daily.**

b) Food contact surfaces are cleaned and sanitized:

1) Before operations begin, food contact surfaces are rinsed with cold water and sanitized with a 100 ppm sodium hypochlorite sanitizer. The sanitation supervisor inspects food contact surfaces to determine if they are clean and have been wetted with sanitizer. **Monitoring Frequency: Daily before operations begin.**

2) During breaks (every 4 hours), major solids are physically removed from floors, equipment, and food contact surfaces. All surfaces are rinsed with cold water. Food contact surfaces are scrubbed, using brushes with a chlorinated alkaline detergent in warm (≥120°F) water. All other surfaces and floors are rinsed with cold water. Food contact surfaces are sanitized with a 100-150 ppm chlorine as sodium hypochlorite sanitizer solution. Floors are sanitized with a 400-600 ppm quaternary ammonium chloride sanitizer. Utensils are cleaned in a deep sink with a warm chlorinated alkaline detergent, rinsed in warm water (≥120°F), and dipped in a 100 ppm chlorine as sodium hypochlorite sanitizer prior to use. The sanitation supervisor checks sanitizers before use and inspects food contact surfaces to determine if they are clean and have been sanitized. **Monitoring Frequency: At the 4- and 8-hour breaks.**

3) At the end of daily operations, major solids are physically removed from floors, equipment, and food contact surfaces. Equipment is disassembled as required for adequate cleaning. All surfaces are rinsed with cold water. Food contact surfaces are scrubbed using brushes with a chlorinated alkaline detergent in warm (≥120°F) water. Floors and walls (splash zone) are washed with warm chlorinated alkaline detergent, using push brooms on floors. All surfaces are thoroughly rinsed with clear water before applying sanitizer. Food contact surfaces are sprayed with 100-150 ppm chlorine as sodium hypochlorite sanitizer solution. Floors and walls are sprayed with a 400-600 ppm quaternary ammonium chloride sanitizer solution. Utensils are cleaned in a deep sink with a chlorinated alkaline cleaner in warm (≥120°F) water, dipped in a 100-150 ppm chlorine as sodium hypochlorite sanitizer and air dried. The sanitation supervisor inspects food contact surfaces to determine if they are clean and have been sanitized. **Monitoring Frequency: Daily at the end of operations.**

c) Workers wear clean gloves and outer garments.

1) Workers working with raw and cooked product wear clean gloves, clean outer garments, and waterproof aprons. Waterproof aprons are cleaned and sanitized twice each day, at the midday break and at the end of the shift.
2) Administrative personnel wear smocks when in processing areas. Smocks are laundered in-house as needed.
3) Maintenance workers wear gray uniforms. Uniforms are laundered in house as needed.
4) Production supervisors monitor the use of gloves and the cleanliness of workers’ outer garments.
   **Monitoring Frequency:** Daily before operations and after each break.

**Corrections:**

a) Food contact surfaces that are not readily cleanable are repaired or replaced.
b) Adjust sanitizer concentration. Food contact surfaces that are not clean are cleaned and sanitized.
c) Gloves that become a potential source of contamination are cleaned and sanitized or replaced. Outer garments that become a potential source of contamination are cleaned and sanitized or replaced.

**Records:**

a-c) Daily Sanitation Control Record (see page 27)

3) **Prevention of cross contamination (FDA Key Sanitation Condition No. 3)**

**Controls and Monitoring:**

a) Production supervisors have received basic food sanitation training. Plant manager schedules basic food sanitation courses for new production supervisors.
   **Monitoring Frequency:** When production supervisors are hired.
b) Employee practices do not result in food contamination (hair restraints, glove use, hand washing, personal belongings storage, eating and drinking, boot sanitizing).

1) Workers wear hairnets, headbands, caps, beard covers, or other effective hair restraints and do not wear jewelry or other objects that might fall into the product, equipment, or containers.
2) Workers wear disposable gloves and replace them as needed.
3) Workers wash their hands and gloves thoroughly and sanitize them before starting work, after each absence from their workstation, and anytime they have become soiled or contaminated.
4) Clothing and personal belongings are not stored in production areas.
5) Workers do not eat food, chew gum, drink beverages, or use tobacco in production areas.
6) Workers wear color-coded aprons (blue in raw product areas and white in cooked product areas) and are not allowed to enter or pass through other processing areas.
7) Workers sanitize their boots in footbaths containing 400 ppm quaternary ammonium chloride sanitizer solution before entering processing areas.
8) Production supervisors monitor employee practices.
   **Monitoring Frequency:** Daily before operations and every 4 hours during production.
c) Boot sanitizing solutions are checked every 4 hours during production. Sanitation supervisor checks boot sanitizing solutions. **Monitoring Frequency: Daily before operations and every 4 hours during production.**

d) Plant grounds are in a condition that protects against contamination of food. Sanitation supervisor inspects plant grounds. **Monitoring Frequency: Daily before operations.**

e) Waste is removed from processing areas during production. Sanitation supervisor monitors removal of waste. **Monitoring Frequency: Daily before operations.**

f) Floors are sloped to facilitate drainage. Processing area floors are inspected for adequate drainage. **Monitoring Frequency: Daily before operations.**

g) Plant buildings are maintained in good repair. Raw-product processing and cooked-product processing areas are separated. Coolers, including the evaporators, are cleaned annually, or more often if needed. Non-food contact surfaces in processing and packaging areas are cleaned daily at the end of the shift. Raw and cooked products are physically separated in coolers. Packaging materials are protected from contamination during storage. Sanitation supervisor inspects plant. **Monitoring Frequency: Daily before operations.**

h) Cleaning and sanitizing equipment is color-coded for specific plant areas: blue for raw-product processing areas, white for cooked-product processing areas, and yellow for toilet facilities and general plant cleaning. Sanitation supervisor observes that proper equipment is used. **Monitoring Frequency: At each cleanup period.**

Corrections:

a) New production supervisors receive basic sanitation instruction.

b) Workers correct deficiencies in hair restraint use, jewelry use, glove use, hand washing, personal belonging storage, eating and drinking in processing areas, and boot sanitizing before working with raw or cooked products.

c) Boot sanitizing solution is changed.

d) Sanitation supervisor initiates correction of potentially contaminating condition.

e) Waste is removed.

f) Floors with standing water will have the drains unplugged, or, if necessary, consultations will be held with plumbing or general contractors and corrections will be made to correct floor drainage problems.

g) Sanitation supervisor initiates correction of potentially contaminating condition, including assessment of product quality.

h) Sanitation equipment that is being used in the wrong plant area is cleaned and sanitized and exchanged for correct equipment. Sanitation supervisor initiates correction of potentially contaminating condition.

Records:

a) Periodic Sanitation Control Record or training record.
b-h) Daily Sanitation Control Record.
4. Maintenance of hand washing, hand sanitizing and toilet facilities (FDA Key Sanitation Condition No. 4)

Controls and Monitoring:

a) Toilet facilities are provided near the workers’ dressing room, physically separated from processing areas. Toilet facilities have self-closing doors, are maintained in good repair, and are cleaned and sanitized daily at the end of operations. Toilet facilities are supplied with toilet paper and other supplies as needed. Sanitation supervisor inspects the toilet facilities and hand washing facilities. **Monitoring Frequency: Daily before operations and every 4 hours during operations.**

b) Handwashing/sanitizing facilities are provided in raw and cooked processing areas and in the toilet facility. Hand washing facilities have: hot and cold running water with foot-activated valves; liquid hand soap; hand sanitizer solutions that are changed every 4 hours during production; sanitary towel service; signs directing workers to wash their hands and gloves thoroughly. Sanitation supervisor inspects the hand washing facilities and checks hand sanitizer strength. **Monitoring Frequency: Daily before operations and every 4 hours during operations.** Note: handwashing procedures are covered under FDA key sanitation condition No. 3, prevention of cross contamination.

Corrections:

a) Sanitation supervisor initiates repairs of toilet or hand washing facilities as needed.

b) Sanitation supervisor restocks facilities or adjusts sanitizers.

Records:

a-b) Daily Sanitation Control Record.

5) Protection from adulterants (FDA Key Sanitation Condition No. 5)

Controls and Monitoring:

a) Cleaning compounds, sanitizers, and lubricants used in processing and packaging areas are approved for use in food plants. Receiving manager checks invoices at receiving before food-grade chemicals are stored. **Monitoring Frequency: When cleaning compounds, sanitizers, and lubricants are received.**

b) Food-grade and non-food-grade chemicals and lubricants are stored separately outside processing and packaging areas. Sanitation supervisor inspects chemical storage areas. **Monitoring Frequency: Daily before operations.**

c) Food, food contact surfaces, and food packaging materials are protected from adulteration from biological, chemical and physical contaminants. Safety-type light fixtures are used in processing and packaging areas. Sanitation supervisor inspects processing and packaging areas. **Monitoring Frequency: Daily before operations and every 4 hours.**

d) Equipment is in good repair with no loose or missing metal parts. Sanitation supervisor inspects processing and packaging equipment. **Monitoring Frequency: Daily before operations.**

e) Drip or condensate does not contaminate food or packaging materials. **Monitoring Frequency: Daily before operations and at 4- and 8-hour breaks.**
Corrections:

a) Unapproved chemicals are returned or used in non-processing areas.
b) Improperly stored chemicals are moved to the correct storage area.
c) Safety of the product is examined and contaminated source is eliminated.
d) Repairs are made as needed to defective equipment.
e) Sanitation supervisor corrects any condensation problems.

Records:

a) Periodic Sanitation Control Record.
b-e) Daily Sanitation Control Record.

6) Labeling, storage and use of toxic compounds (FDA Key Sanitation Condition No. 6)

Controls and Monitoring:

a) All toxic compounds used in the plant are labeled with the manufacturer’s name, use instructions, and the appropriate EPA approval, or have documentation with the necessary information. Receiving manager verifies that this information is present before toxic compounds are stored. **Monitoring Frequency: When toxic compounds are received.**
b) Cleaning compounds, sanitizing agents, lubricants, pesticide chemicals, and other toxic compounds are properly labeled and stored in a closed and locked cage in dry storage outside processing and packaging areas and separately from food-grade chemical, food-grade lubricant, and packaging material storage. Only authorized personnel have access to the cage. Sanitation supervisor checks cage for cleanliness and container leakage. **Monitoring Frequency: Daily before operations.**
c) All manufacturers’ instructions and recommendations are followed. Only authorized personnel fill small working containers, such as containers of hand sanitizing compounds. These containers are properly marked with the common name of the chemical and are not stored in any way that may cause the chemical to fall or drip onto food or food-packaging materials. Sanitation supervisor verifies proper procedures and labeling. **Monitoring Frequency: Daily before operations.**

Corrections:

a) Toxic compounds without proper information are placed on hold until information is obtained. Toxic compounds without documentation are returned to the supplier.
b) Improperly stored chemicals are moved to the correct storage area. Leaking containers are resealed or replaced as necessary. Storage cage will be cleaned by the next working day.
c) Misuse of toxic compounds results in disciplinary action or retraining. Potentially contaminated food is discarded or destroyed. Improper labeling of working containers is corrected.

Records:

a) Periodic Sanitation Control Record.
b-c) Daily Sanitation Control Record.
7) Employee health (FDA Key Sanitation Condition No. 7)

Controls and Monitoring:

a) Workers report to their immediate supervisor any health condition that might result in food contamination. Supervisors report suspected health problems to the plant manager. The plant manager decides if a potential food contamination situation exists. **Monitoring Frequency: Daily before operations.**
b) Supervisors check for infected lesions that might contaminate food. **Monitoring Frequency: Daily before operations.**

Corrections:

a) Workers who represent a potential risk are sent home or reassigned to non-food contact jobs.
b) Cover lesion with impermeable bandage, reassign, or send worker home.

Records:

a-b) Daily Sanitation Control Record.

8) Exclusion of pests (FDA Key Sanitation Condition No. 8)

Controls and Monitoring:

a) A pest management firm treats the outside of the building. They also inspect the interior of the building and treat as necessary with appropriate chemicals. **Monitoring Frequency: Every other month.**
b) Plant grounds and interior areas are kept free of litter, waste, and other conditions that might attract pests. Outer plant doors are kept closed, processing areas are screened with plastic curtains, and electric bug-killing devices are located outside entrances to processing areas. No pets are allowed in the plant. Supervisors report any pest problems to the plant manager. The sanitation supervisor inspects for the presence of pests. **Monitoring Frequency: Daily before operations.**

Corrections:

a) Conditions that may cause pest problems are corrected.
b) The pest management firm is notified of any pest problem and treats the problem. Pest treatments are more frequent if problems are identified.

Records:

a) Periodic Sanitation Control Record.
b) Daily Sanitation Control Record.
### Table 2

Seafood HACCP Regulation Sanitation Requirements (21 CFR 123.11(b)) and their relation to the current Good Manufacturing Practice Regulation (21 CFR 110)

<table>
<thead>
<tr>
<th>Part 123.11(b) Monitoring Requirement</th>
<th>Corresponding Part 110 Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Safety of water</td>
<td>.37(a): Process water safe and of adequate sanitary quality; water used for ice manufacture of safe and adequate sanitary quality&lt;br&gt;.37(b)(5): No cross connections between sewer or wastewater and process water</td>
</tr>
<tr>
<td>Condition and cleanliness of food contact surfaces</td>
<td>.40(a)(b): Food contact surfaces designed, fabricated, maintained, and installed to be adequately cleanable and to withstand the environment of use and cleaning compounds; smoothly bonded seams&lt;br&gt;.35(d)(2) &amp; .80(b)(1), (b)(10), (b)(13)(ii): When cleaning is necessary to protect against introduction of microorganisms, clean and sanitize before use, after interruptions, and as necessary&lt;br&gt;.10(b)(1) &amp; (5): Gloves should be impermeable, clean, and sanitary; outer garments suitable</td>
</tr>
<tr>
<td>Prevention of cross-contamination</td>
<td>.10(b), (b)(2), (b)(3), (b)(4), (b)(7), (b)(8), (b)(9) &amp; .80(b)(6), (b)(13)(iv): Food handlers conform to hygienic practices to the extent necessary to prevent contamination; maintain adequate personal cleanliness; wash, and sanitize if necessary, hands before start work, after absence from work station, and when become contaminated; taking precautions as necessary to protect against contamination with microorganisms; effective measures to prevent finished product contamination by raw materials, other ingredients, or refuse; remove jewelry that cannot be sanitized; abstaining from eating, chewing gum, drinking, or using tobacco near exposed food or equipment; storing clothing or personal items away from exposed food and equipment&lt;br&gt;.20(b)(1), (2), (4): Plant design must reduce potential for contamination of food, food contact surfaces, and packaging material and must permit employees to protect against contamination of food from clothing or personal contact; separation of operations.</td>
</tr>
<tr>
<td>Maintenance of hand washing, hand sanitizing, and toilet facilities</td>
<td>.37(e), (e)(1), (e)(2), (e)(3), (e)(4): Hand washing and, where appropriate, hand sanitizing facilities should be at each location where good sanitary practice dictates their use; effective hand cleaning and sanitizing preparations; water at suitable temperature; sanitary towel service or suitable drying devices; designed to prevent recontamination&lt;br&gt;.37(c) &amp; (d): Adequate sewage disposal system; adequate, readily accessible toilet facilities; maintained in sanitary condition; self closing doors; protect food from airborne contamination.</td>
</tr>
<tr>
<td>Protection from adulterants</td>
<td>.40(a), .80 &amp; .80(a)(5), (a)(7) &amp; (b)(5), (b)(7), (b)(10), (b)(12), (b)(13) &amp; 93: Design, construction, and use of equipment precludes adulteration of food with lubricants, fuel, metal fragments, contaminated water, or other contaminants; all reasonable measure to ensure that production methods do not contribute contamination; raw materials held to protect against contamination; work-in-progress handled to protect against contamination; equipment protects food from contamination; mechanical steps protect food from contamination; batters, breading, sauces, dressings, etc. protected from contamination; filling, assembly, packaging, and other operations protect food from contamination; storage and transportation protect the food from contamination.&lt;br&gt;.20(b)(4) &amp; .80(b)(10), (b)(12)(iv): Drip or condensate from fixtures, ducts and pipes does not contaminate food, food contact surfaces, or packaging material; adequate physical protection of food from contaminants that may drip, drain, or be drawn into the food should be provided.&lt;br&gt;.40(lg): Compressed air or other gases mechanically introduced treated to prevent contamination of food</td>
</tr>
<tr>
<td>Proper labeling, storage and use of toxic compounds</td>
<td>.35(b)(2) &amp; (c): Toxic cleaning compounds, sanitizing agents, and pesticides identified, held, and stored in a manner that protects food, food contact surfaces, and packaging material from contamination; all relevant regulations for their use followed; pesticides used only when food, food contact surfaces, and packaging material protected from contamination</td>
</tr>
<tr>
<td>Control of employee health conditions</td>
<td>.10(a): Food handler who has illness or open lesion or other source of microbiological contamination that presents reasonable possibility of contamination of food, food contact surface, or packaging material excluded from such operations</td>
</tr>
<tr>
<td>Exclusion of pests</td>
<td>.35(c): No pests shall be allowed in any area of a food plant</td>
</tr>
</tbody>
</table>
### Daily Sanitation Control Record

**Report Date:**

<table>
<thead>
<tr>
<th>Sanitation Area and Goal</th>
<th>Pre-Op Time</th>
<th>Start Time</th>
<th>4 Hour Time</th>
<th>8 Hour Time</th>
<th>Post-Op Time</th>
<th>Comments and Corrections</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Safety of water</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>(See Monthly Sanitation Control Record)</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Back Siphonage – Hose (S/U)*</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2) Condition and cleanliness of food contact surfaces</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>(See Monthly Sanitation Control Record)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Equipment cleaned and sanitized</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Line 1: (S/U)</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Line 2: (S/U)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>• Sanitizer Strength</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Sanitizer Type</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Strength</td>
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<tr>
<td>ppm</td>
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<tr>
<td>Line 1: (ppm)</td>
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<tr>
<td>Line 2: (ppm)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Gloves and aprons clean and in good repair</td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>Line 1: (S/U)</td>
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<td>Line 2: (S/U)</td>
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<td></td>
</tr>
<tr>
<td>3) Prevention of cross-contamination</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>(See Monthly Sanitation Control Record)</td>
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<td></td>
</tr>
<tr>
<td>• Hands, gloves, equipment, and utensils washed/sanitized after contact with unsanitary objects (S/U)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Employees working on raw products, wash and sanitize hands/ gloves/outerwear before working with cooked products (S/U)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Unpackaged cooked products separated from raw products (S/U)</td>
<td></td>
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</tr>
</tbody>
</table>

*S = Satisfactory / U = Unsatisfactory
<table>
<thead>
<tr>
<th>Sanitation Area and Goal</th>
<th>Pre-Op Time</th>
<th>Start Time</th>
<th>4 Hour Time</th>
<th>8 Hour Time</th>
<th>Post-Op Time</th>
<th>Comments and Corrections</th>
</tr>
</thead>
<tbody>
<tr>
<td>4) Maintenance of hand washing, hand sanitizing, and toilet facilities</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Hand washing and hand sanitizing stations adequate</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Line 1: (S/U)</td>
<td></td>
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</tr>
<tr>
<td>Line 2: (S/U)</td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>• Hand sanitizing station</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Sanitizer Type</td>
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</tr>
<tr>
<td>Strength ppm</td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Line 2: (ppm)</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>• Toilets clean, properly functioning, and adequately supplied (S/U)</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>5) Protection from adulterants and 6) Labeling, storage, and use of toxic compounds</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Product protected from contamination (S/U)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Cleaning compounds, lubricants, and pesticides labeled and stored properly (S/U)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7) Employee health</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>• Employees do not show signs of medical problems (S/U)</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>8) Exclusion of pests</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Pests excluded from processing area (S/U)</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>S = Satisfactory / U = Unsatisfactory</td>
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</tr>
<tr>
<td>Signature or initials __________________________________</td>
<td>Date ____________________________</td>
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</tr>
</tbody>
</table>
### Monthly Sanitation Control Record

<table>
<thead>
<tr>
<th>Sanitation Area</th>
<th>Decision</th>
<th>Comments/Corrections</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Safety of water</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Safe and sanitary source (S/U) (Annual)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• No cross-connections in hard plumbing (S/U)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2) Condition and cleanliness of food contact surfaces</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Processing equipment and utensils in suitable condition (S/U)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3) Prevention of cross-contamination</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Physical conditions of plant and layout equipment (S/U)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

S = Satisfactory / U = Unsatisfactory

Additional Comments:

Signature or initials: ____________________________________________________________
### Periodic Sanitation Control Record

**Periodic Sanitation Control Record**

<table>
<thead>
<tr>
<th>Condition</th>
<th>S</th>
<th>U</th>
<th>Comments/Corrections</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Safety of water:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a) Municipal water bill (annually).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>b) Building plumbing inspection report</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(when plumbing is modified)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3) Prevention of cross-contamination:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a) Production supervisors have received</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>basic food sanitation training (when</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>hired).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5) Protection from adulteration:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a) Invoices for food-grade chemicals</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>checked at receiving before chemicals</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>are stored.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6) Labeling, storage and use of toxic</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>compounds:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a) Labels or documents for toxic compounds</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>checked at receiving before compounds are</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>stored.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8) Exclusion of pests:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a) Pest management firm’s report is satisfactory (every other month).</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Firm Name:** ____________________________  **Date:** ________________

**Firm Address:** __________________________

**Condition S U Comments/Corrections**

- S = Satisfactory / U = Unsatisfactory

**Comments and Corrections:**

Report by: ____________________________
Preliminary Steps in Developing a HACCP Plan

Before building a HACCP plan, there are preliminary steps that should be considered (Slide 1).

1) Assemble HACCP Team

Assembling a HACCP team is an important step in building a HACCP plan. The team should consist of individuals with different specialties and experience with the process. The HACCP team should include members who are directly involved with the plant’s daily operations. The team may include personnel from maintenance, production, sanitation, quality control and laboratory. They develop the HACCP plan, write SCPs, and verify and implement the HACCP
system. The team members should be knowledgeable about food safety hazards and HACCP principles. When issues arise that cannot be resolved internally, it may be necessary to enlist outside expertise.

In addition to writing and developing the HACCP plan, the HACCP team provides oversight of the implementation of the plan into the daily operations of the facility. This includes ensuring that applicable personnel are trained appropriately to handle their required duties.

Although one person may be able to analyze hazards and develop a HACCP plan successfully, many companies find it helpful to build a HACCP team. When only one person develops the HACCP plan, some key points can be missed or misunderstood in the process. The team approach minimizes the risk that key points will be missed or that aspects of the operation will be misunderstood. It also encourages ownership of the plan, builds company involvement and brings in different areas of expertise.

In small companies, the responsibility for writing the HACCP plan may fall to one person. If it is possible to build a HACCP team in a small company, employees knowledgeable of various divisions, including owners, should be members of the HACCP team. Universities, Cooperative Extension, consulting groups, Sea Grant programs, trade associations, model plans and published guidance can provide additional assistance (see chapter 13 for more information).

2) Describe the Product, Intended Use and Consumers

The HACCP team should describe the product(s), the type of packaging, the method of distribution, the intended customer (e.g., general public, infants, elderly) and likely use of the product (e.g., consumed without further cooking, heat-and-serve, cooked). It may seem to take a lot of effort to complete a very detailed description of the product and intended use, but it is necessary to assure an accurate hazard analysis (Slide 3).

Slide 3

Product Description should include:
- Type of seafood product (species and finished product form)
- Where product is purchased
- How product is received, stored, and shipped
- How product is packaged
- Intended end use

A complete product description should include:

Type of seafood:
- species of fish or shellfish including market name (e.g., tuna) and scientific name (e.g., Thunnus albacares), if necessary
- finished product form (e.g., raw, cooked, pasteurized, smoked, etc.)
It is important to know which species are being processed in order to accurately identify potential food safety hazards. For the same reason, it is important to know the finished product form.

Identify where product is purchased:

- directly from the fisherman;
- directly from the grower/farmer;
- from another processor; or
- from a combination of these sources.

It is important to know where the product is purchased to identify the correct potential food safety hazards. For example, how a potential hazard is addressed will depend on whether the processor is a “primary processor” or a “secondary processor.” A “primary processor” is the first receiver of the product from a harvest vessel or aquaculture site. A “secondary processor” receives product from a “primary processor” or another “secondary processor.”

Identify how the fish are received:

- fresh – under refrigeration;
- fresh – under ice or chemical coolant;
- frozen;
- canned or shelf-stable;
- more than one of these methods.

Identify how the fish are stored after receipt:

- fresh – under refrigeration;
- frozen;
- dry storage.

Identify how the finished product will be shipped:

- fresh – under refrigeration;
- fresh – under ice or chemical coolant;
- frozen;
- ambient conditions;
- more than one of these methods.

Identify how the finished product will be packaged:

- air-permeable packaging (e.g., foam tray overwrapped with fresh meat film);
- reduced-oxygen packaging (e.g., vacuum packaging, modified-atmosphere packaging, controlled-atmosphere packaging, hermetically-sealed packages, or packed in oil).

It is important to know how products are received, stored, packaged, and shipped to identify the correct potential food safety hazards.
Identify how the products are intended to be consumed:

- to be cooked by the consumer;
- ready-to-eat (RTE) raw;
- RTE – cooked
- RTE – partially cooked;
- RTE – heat and serve;
- RTE – reheat.

It is important to know how the product will be consumed to identify any potential food safety hazards.

A Product Description form (Slide 4) has been developed to help record this information.

---

### Slide 4

**Product Description Form for Fish and Shellfish Species**

<table>
<thead>
<tr>
<th>Type of Seafood Product (Species name)</th>
<th>Where Product Is Purchased (Source)</th>
<th>How Product Is Received</th>
<th>How Product Is Stored</th>
<th>How Product Is Shipped</th>
<th>How Product Is Packaged</th>
<th>Intended Use</th>
<th>Intended Consumer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>From Fisherman</td>
<td>Refrigerated</td>
<td>Frozen</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>General Public</td>
<td>At-risk Population</td>
</tr>
<tr>
<td></td>
<td>From Fish Farm</td>
<td>Refrigerated</td>
<td>Frozen</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>General Public</td>
<td>At-risk Population</td>
</tr>
<tr>
<td></td>
<td>From Processor</td>
<td>Refrigerated</td>
<td>Frozen</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>General Public</td>
<td>At-risk Population</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Refrigerated</td>
<td>Frozen</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>General Public</td>
<td>At-risk Population</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Refrigerated</td>
<td>Frozen</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>General Public</td>
<td>At-risk Population</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Refrigerated</td>
<td>Frozen</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>General Public</td>
<td>At-risk Population</td>
</tr>
</tbody>
</table>

---

### 3) Develop an Accurate Process Flow Chart

A flow chart provides an important visual tool that the HACCP team can use to identify and describe the process (Slide 5). When developing a process flow chart, it is important to include all the process steps within the facility’s control from receiving through final product storage, including reworked product if applicable. Since the accuracy of the process flow is critical to conduct a hazard analysis, the steps outlined in the chart must be verified at the plant. If a step is missed, a significant food safety hazard may be missed. Include every handling,
processing and holding step for primary product as well as ingredients and packaging. The HACCP team should walk through the facility and make any changes required in the flow chart. The walk-through allows each team member to gain an overall picture of how the product is made. It may be helpful to invite additional plant personnel to review the diagram during the walk-through.

4) Describe the Process

A written process description can be a useful tool to explain what happens at each of the process steps needed to produce a product covered by a particular HACCP plan. This description can be used as a working reference for the development of the HACCP plan and to facilitate communication with company personnel and regulators. It is also important to know what occurs at each process step. For example, it is important to know information about process steps such as the maximum length of time that the product could be exposed to unrefrigerated temperatures, the maximum room air temperature, or the maximum internal product temperature. This information is necessary to conduct an accurate hazard analysis.

Example: XYZ Seafood Co.

The model XYZ Seafood Company is used to illustrate the progressive development of the HACCP program beginning with the three preliminary steps through the application of each of the HACCP principles.

1) Assemble HACCP Team

XYZ Seafood Company has determined that their team will consist of three individuals including: the plant manager, production supervisor and the sanitation supervisor. All have undergone seafood HACCP training.

2) Describe the Product, Intended Use and Consumers

XYZ Seafood Company used the Product Description Form for Fish and Shellfish Species (Slide 4) to complete this preliminary step. Results are provided in Slide 6.
The results from this chart are summarized below:

**Product Description:** Raw, wild-caught mahi-mahi fillets

**Fishery Product Market Name:** Mahi-mahi (*Coryphaena* species)

**Source of Fishery Product:** From other processors, received in ice

**Methods of Packaging, Distribution and Storage:** Air packed, stored and distributed on ice

**Intended Use and Consumer:** To be cooked and consumed by the general public

3) **Develop a Process Flow Chart**

XYZ Seafood Company developed a process flow chart (Slide 7).
4) Develop a Process Description

XYZ Seafood Company’s HACCP team developed a written description for each of the steps in their process flow chart.

**Receive fresh fish** – Fresh wild caught mahi-mahi (*Coryphaena* species, not aquacultured) fillets are received from several domestic suppliers. Delivery truck transit times range from 2 to 8 hours. Tubs or other containers of mahi-mahi fillets are received along with other fresh seafood products packed in ice and delivered by refrigerated truck. After receipt, products are re-iced if necessary and moved into refrigerated storage.

**Refrigerated storage** – Individual mahi-mahi fillets are completely buried in ice and stored in a refrigerated cooler until needed.

**Trim** – Individual tubs or containers of mahi-mahi fillets are removed from the cooler as needed to pack customer orders. Fillets are trimmed by hand with knives if necessary to meet customer specifications. Trimming is completed in 30 minutes or less.

**Weigh/Pack/Label** – Per customer order, mahi-mahi fillets are weighed, packed into containers, and each container is labeled with a handwritten or printed label that contains the market name of the species of fish that it contains. Individual containers are completely surrounded by ice and assembled into master cartons for each customer order. The weigh/pack/label steps are completed in 30 minutes or less.

**Finished product storage** – Iced mahi-mahi containers in master cartons that contain each customer’s order are placed back into refrigerated storage until it is moved directly to refrigerated trucks for delivery to retail or restaurant customers.
It is important in developing a HACCP plan or modifying an existing plan to be aware of the potential seafood safety hazards that are associated with the products and processes. The rationale for implementing a seafood HACCP program is to prevent the potential seafood safety hazards that are “reasonably likely to occur” and could cause disease or injury if not adequately controlled (Slide 1). Awareness of the potential hazards must also include some knowledge about the most appropriate and effective controls. This chapter provides a brief introduction to some of the most common hazards of concern. The FDA's Fish and Fisheries Products Hazards and Controls Guidance (Hazards Guide), provides step-by-step directions to assist in determining the specific hazards and their respective controls. Controls can differ by product types and processes. Use of the Hazards Guide will be explained in Chapter 5-Hazard Analysis.

In this chapter you will learn about:
• Food Safety Hazards that have been associated with seafood and are considered “reasonably likely to occur” if not subject to appropriate controls.

In developing a HACCP plan, it is important to understand that seafood hazards only refer to the conditions or contaminants in food that can cause illness or injury to people (Slide 2). Many conditions are highly undesirable in food, such as the presence of insects, hair, filth or spoilage. Economic fraud and violations of regulatory food standards are equally undesirable. Processors should have prerequisite programs in place to properly address all of these conditions. However, since these conditions are often not directly related to the safety
of the product, it is recommended that they **should not** be included within a HACCP plan.

**Slide 2**

**Hazards:** a biological, chemical or physical agent that is reasonably likely to cause illness or injury in the absence of appropriate controls.

**Undesirable conditions** may not impose a particular food safety hazard, but they are subject to other regulatory controls and pre-requisite requirements (i.e., GMPs and Sanitation Control Procedures (SCPs)). Examples include:
- Insects
- Hair
- Filth
- Spoilage
- Economic fraud
- Violations of regulatory food standards not directly related to safety

The variety of potential seafood safety hazards can be grouped into three categories (Slide 3):

**Slide 3**

Potential seafood safety hazards can be grouped in three categories:
- Biological Hazards
- Chemical Hazards
- Physical Hazards

- **Biological hazards** including harmful bacteria, viruses or parasites.
- **Chemical hazards** including compounds or ingredients that can cause illness due to immediate or long-term exposure.
- **Physical hazards** including presence of foreign objects, such as glass or metal fragments.

It is not within the scope of this course to go into detail on the seafood hazards and respective controls. This topic is too large and would be covered better in separate microbiology, toxicology and food-processing courses. The purpose of this chapter is to help participants develop an awareness of the kinds of hazards that may occur in seafood and to learn some of the general controls to prevent these hazards. Food processors may find it necessary to work with technical experts with more knowledge and experience with potential seafood safety hazards for various products and processes.

**Biological Hazards**

All foods can contain biological hazards. These hazards can come from raw materials or from food-processing steps used to make the final product. The
biological hazards typically involve living organisms too small to be seen with the naked eye. They are called by the generic term “microorganisms.” Microorganisms are classified into various groups. A few groups important in foods include: yeasts, molds, bacteria, viruses and parasites (Slide 4). Table 1 at the end of the chapter lists the types of materials that can be biological hazards in foods.

Microorganisms are found everywhere: air, dirt, fresh and salt water, skin, hair, animals and plants. People may come into contact with thousands of kinds of yeasts, molds, bacteria, viruses and parasites daily without ill effect. Since microorganisms are so widespread, it is important to understand when to be concerned about them and how to control them if necessary. It is important to understand the different types of microorganisms in order to distinguish the types that cause foodborne illnesses.

Many microorganisms are beneficial. Certain kinds of yeast, molds and bacteria help make cheese, sour cream, yogurt and other fermented dairy products. Certain kinds of yeast are used in making beer, wine and other fermented beverages. We add these microorganisms to our foods intentionally, and they cause no harm. In fact, studies show that some of these microorganisms contribute to good health. The term “probiotics” is being used for live microorganisms placed in food to provide a health benefit to the consumer.

Although thousands of different microorganisms exist, only a few pose hazards to humans. These hazardous microorganisms are called pathogens. Pathogens that are transmitted by food are called foodborne pathogens or seafoodborne pathogens. Among the five groups of microorganisms described earlier, only bacteria, viruses and parasites are typically included among those that can make seafood unsafe (Slide 5). Generally, yeast and molds do not pose a biological hazard in seafood. Some molds produce hazardous toxins, but these toxins are considered chemical hazards.

Microorganisms that can be pathogenic and cause seafoodborne illnesses:
• Bacteria
• Viruses
• Parasites
Bacterial Hazards – Infections and Intoxication

Bacterial hazards are defined as those bacteria that, if they occur in food, may cause illness in humans, either by infection or intoxication (Slide 6). Foodborne infections are caused by consuming live pathogens that grow within the body, usually in the intestinal tract. Foodborne infections can be visualized much the same as an infection on your skin, except that it is an infection on the surface of the intestinal tract. Foodborne infections are caused by organisms such as Salmonella spp., Shigella spp., and Listeria monocytogenes to mention a few. If certain types and amounts of these particular bacteria are simply present in the seafood when consumed, they can grow and infect the consumer.

The other kind of bacterial hazard is foodborne intoxication, which is a condition caused by swallowing preformed toxins (i.e., toxins produced by microorganisms in the food before it is eaten). The onset of symptoms from a foodborne intoxication is often more rapid than symptoms from a foodborne infection that requires more time for the bacteria to grow after consumption. Evidence of an illness due to seafoodborne intoxications can occur within minutes to hours after consumption, whereas most foodborne infections are not evident until hours or days after consumption. Foodborne intoxications can be caused by certain types of bacteria such as Staphylococcus aureus or Clostridium botulinum that can produce staph enterotoxins or botulinum neurotoxins, respectively. If certain types of these bacteria are allowed to grow in the seafood prior to consumption, they can produce toxins to levels that could cause intoxication of the consumer.

Bacterial Hazards – Presence, Growth and Survival

Certain bacteria can be a potential hazard simply due to presence in or on the seafood in any amount when consumed, whereas other types may require growth to a level or amount that is more hazardous. Knowledge regarding bacterial hazards due to presence or growth is important relative to selection of appropriate controls. Based on the nature of seafood harvests and aquaculture farming, it is reasonable to assume that some potentially hazardous bacteria will be present on live and raw products, but cooking can be a very effective control. An effective cooking procedure can significantly reduce or eliminate the pathogens of concern. Improper cooking could allow the pathogens to survive in amounts that still pose a hazard. Selection of appropriate controls must always consider potential bacteria survival.

In some cases, seafood is eaten raw or as ready-to-eat (RTE) products that do not require cooking just before consumption (Slide 7). These products would require additional controls to reduce or eliminate certain types and amounts of
Seafood Safety Hazards

bacteria that may be present at harvest or introduced as contaminants during processing and handling.

Slide 7

- **Seafood products commonly eaten raw**: oysters, clams, conch and sushi
- **Seafood Ready-To-Eat (RTE) products**: pre-cooked and frozen shrimp, smoked fish, pickled fish, pasteurized crab meat, and pasteurized surimi (fabricated seafood analog)

Bacterial growth occurs in phases beginning with a slow or lag phase that provides time for adjustment to the growing conditions (Slide 8). Eventually under favorable conditions, the bacteria growth can rapidly accelerate with exponential growth or doubling as each bacterium grows large and divides. One bacterium divides into two, two into four, four into eight, eight into sixteen, and so on. Under ideal conditions, some bacteria double every 20 minutes. One bacterium can multiply to more than 30,000 bacteria in five hours and over 16 million in eight hours.

Slide 8

<table>
<thead>
<tr>
<th>Growth Time</th>
<th>Lag Phase</th>
<th>Exponential Phase</th>
<th>Stationary Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Bacteria</td>
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</tr>
</tbody>
</table>

Effective controls remove or reduce the favorable growth conditions (Slide 9). Bacteria prefer favorable temperatures, moisture and certain atmospheric conditions (air or reduced-oxygen atmospheres) for growth. Hazard prevention usually involves controls and monitoring of these conditions to restrict and eliminate bacterial growth.
Bacterial Hazards – Sporeformers and Non-Sporeformers

Bacterial hazards can be grouped into sporeformers and non-sporeformers. Certain types of bacteria (e.g., *Clostridium* spp. and *Bacillus* spp.) pass through a dormant stage in their life cycle called a spore. When a bacterium exists as a spore, it is very resistant to chemicals, heat and other treatments that would normally be lethal to non-sporeforming bacteria. Because they are dormant, spores are not hazardous as long as they stay spores. Unfortunately, if they survive a processing step designed to kill non-sporeforming bacteria, they may become a hazard in the food if they are allowed to germinate from the spore form and grow. When sporeformers are a concern, the process steps used to control them are often much more rigorous than necessary to control non-sporeformers (Slide 10).

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The following list of potential foodborne bacterial pathogens includes those that are more “reasonably likely to occur” in seafood, but this listing is not fully inclusive. Additional, emerging pathogens may be added over time based on experience and discovery.

*Clostridium botulinum* (Sporeformer)

*Clostridium botulinum* is found throughout the environment and has been isolated from soil, water, vegetables, meats, dairy products, ocean sediments, the intestinal tracts of fish, and the gills and viscera of crabs and other shellfish.
C. botulinum is a sporeforming bacterium that grows in the absence of air. These characteristics allow it to survive normal cooking temperatures and to grow in a vacuum-packaged and modified-atmosphere environment. C. botulinum produces a powerful neurotoxin that causes botulism. Growth is necessary for C. botulinum to produce toxin. Symptoms include diarrhea, vomiting, abdominal pain, nausea and weakness. These are followed by double, blurred vision and dilated, fixed pupils. In severe cases, paralysis of the muscles responsible for breathing can cause death.

There are various types of C. botulinum (Type A, B, C, D, E, F and G) that are classified according to the type of neurotoxin they can produce. C. botulinum Type E is the most common type associated with fish and fishery products. It is a particular concern because it grows at temperatures as low as 38°F and can produce little noticeable evidence of spoilage. In contrast, C. botulinum Type A is the form of this bacteria that is most common in land-based products (i.e., vegetables). It is a common contaminant on processing equipment. It will grow at temperatures above 50°F and produces a putrid odor in products in which it grows. However, its spores are much more heat-resistant than the Type E form of the bacteria. Thus controls for Type E and Type A would differ.

Because C. botulinum produces heat-resistant spores and requires the absence of oxygen for growth, botulism has been most commonly associated with improperly canned food (usually home canned). Semi-preserved seafood, including smoked, salted and fermented fish, has also been identified as a cause of botulism. Packaging of raw, non-frozen fishery products in reduced-oxygen or with restricted-oxygen exposure (i.e., anaerobic packaging such as vacuum packs with oxygen impermeable film) is also considered favorable for potential growth of C. botulinum Type E if the packages are not properly refrigerated.

C. botulinum Type E can be controlled by inhibiting growth of the bacteria or by destroying it in the seafood (Slide 11). Proper thermal processes for canned seafood destroy the bacteria. Heavy salting or drying to reduce the water activity (a) below 0.93 and fermentation or acidification to below pH 4.6 are effective means of preventing C. botulinum growth. Maintaining proper storage temperatures alone is not considered an adequate control measure for C. botulinum Type E because of its ability to grow at low temperatures and because of the severity of the illness. In many products, it is an important second barrier to growth. Additional controls are necessary for raw, non-frozen seafood packaged in anaerobic (limited oxygen) conditions that could allow spore germination.

**Bacillus cereus (Sporeformer)**

Bacillus cereus spores can germinate and grow in the presence or absence of oxygen. If allowed to grow, it can produce heat stable Bacillus toxins that cannot be destroyed by routine cooking. They occur in soils and can be easily transmitted to many types of foods. They have been associated with illnesses involving cooked vegetables and meats, rice, and dairy products. They are not commonly associated with popular seafood products, but their occurrence in foodborne illnesses is considered somewhat underreported. Difficulty in detection and reporting is partially due to the variation in symptoms that can
appear similar to other pathogenic bacteria, and the symptoms are relatively mild and of short duration. *B. cereus* can cause two types of illnesses. The more common is diarrhea and abdominal pains that can occur within 4 to 16 hours after consumption of contaminated food, followed by symptoms for up to 24 hours. The less frequent illnesses involve a more acute onset of nausea and vomiting within 1 to 5 hours which can persist for up to 24 hours after consumption.

Controls are necessary to prevent temperature abuse and prolonged storage or display of previously prepared foods without refrigeration (Slide 12). A primary concern is with precooked, chilled food products with expected long shelf lives. It is important to use rapid chilling methods to reduce the temperature of hot prepared foods like soups. It is best to cool hot foods below 41ºF (7ºC) within less than 4 hours after preparation, display or serving. Likewise, it is important to reheat cold prepared foods rapidly to above 165ºF (74ºC).

**Slide 11**

Some controls for *Clostridium botulinum* in seafood:
- Destroy spores during processing (e.g., thermal processing (canning) or proper cooking to destroy the spores).
- Prevent potential growth by proper salting, drying, or pickling (acidification).
- Proper refrigeration, particularly for raw, non-frozen seafood packaged in anaerobic conditions (limited oxygen).
- Packaging refrigerated fishery products in permeable film that allows enough oxygen exposure to prevent anaerobic growth.

**Slide 12**

Some controls for *Bacillus cereus* in seafood:
- Proper sanitation to prevent product contamination (product source, process facilities and personnel)
- Proper chilling rates for warm prepared food
- Proper refrigeration for prepared, ready-to-eat (RTE) food with extended shelf lives

**Listeria monocytogenes** (Non-sporeformer)

*Listeria monocytogenes* is widespread in nature and has been isolated from soil, vegetation, marine sediments and water. In the early 1900s, *L. monocytogenes* was recognized as a bacterium that caused illness in farm animals. More recently, it has been identified as the cause of listeriosis in humans. Most healthy individuals are either unaffected by *L. monocytogenes* or experience only mild flu-like symptoms. Victims of severe listeriosis are usually immunocompromised. Those at highest risk include cancer patients, individuals taking drugs that affect the body’s immune system, alcoholics, pregnant
women, persons with low stomach acidity, and individuals with AIDS. Severe listeriosis can cause meningitis, abortions, septicemia and a number of other maladies, some of which may lead to death.

The greatest threat of listeriosis is from ready-to-eat products that do not require further cooking before they are eaten (RTE). The *L. monocytogenes* in raw food that will be cooked before consumption is less of a concern to the food industry since the bacteria are killed during cooking. *L. monocytogenes* has been isolated from raw fish, cooked crabs, raw and cooked shrimp, raw lobster, smoked fish and surimi (seafood analog). One of its most significant characteristics is its ability to grow at temperatures as low as 31°F.

*L. monocytogenes* can be controlled by thorough cooking of the seafood and by preventing cross contamination once the seafood is cooked (Slide 13). Since the infective dose of *L. monocytogenes* is thought to be small, time/temperature abuse of food products may not be necessary to cause illness. As an additional safeguard, temperature controls for ready-to-eat seafood is recommended to control potential growth.

**Slide 13**

Some controls for *Listeria monocytogenes* in seafood:
- Proper sanitation to prevent product contamination (product source, process facilities, and personnel)
- Proper refrigeration to prevent growth
- Proper cooking
- Prevent cross-contamination after cooking

**Salmonella spp. (Non-sporeformer)**

*Salmonella* is naturally found in the intestinal tracts of mammals, birds, amphibians, and reptiles, but not in fish, crustaceans or mollusks. *Salmonella* is transferred to seafood through sewage pollution of the harvest environment or by contamination after harvest.

*Salmonella* food infection causes nausea, vomiting, abdominal cramps and fever. Outbreaks of *Salmonella* food infection have been associated with raw oysters, salmon, tuna salad, shrimp cocktail, stuffed sole, and gefilte fish.

*Salmonella* can be prevented by heating seafood sufficiently to kill the bacteria, holding chilled seafood below 40°F, preventing cross-contamination after cooking, and prohibiting people who are ill or are carriers of *Salmonella* from working in food operations (Slide 14). The infective dose of *Salmonella* is thought to be extremely variable, relatively high for healthy individuals and very low for at-risk individuals, such as the elderly or medically compromised. For this reason, illness could result even without time/temperature abuse, but temperature abuse has been a contributing factor in many outbreaks.
**Staphylococcus aureus** (Non-sporeformer)

Humans and animals are the primary reservoirs or source for *S. aureus*. Natural sources include the nose, throat and on the hair and skin of healthy individuals. However, the bacteria can also be found in air, dust, sewage and surfaces of food processing equipment. *S. aureus* can produce a toxin if allowed to grow in food. The toxin is not destroyed by cooking or other thermal process (i.e. canning processes). *S. aureus* has the ability to grow and produce toxins in food with very little available water (0.85 a_w, 10 percent salt), which would prevent the growth of other pathogens.

*Staphylococcus aureus* food poisoning causes nausea, vomiting, abdominal cramping, watery or bloody diarrhea, and fever.

*S. aureus* can be controlled by minimizing time/temperature abuse of seafood, especially after cooking, and requiring that food handlers practice proper hygiene (Slide 15).

**Slide 14**

Some controls for *Salmonella* spp. in seafood:
- Proper sanitation to prevent product contamination (product source, process facilities and personnel)
- Proper refrigeration to prevent growth
- Proper cooking
- Prevent cross-contamination after cooking

**Vibrio cholerae** (Non-sporeformer)

*Vibrio cholerae* is found in estuaries, bays, and brackish waters. It is naturally occurring and is not necessarily related to sewage contamination. *V. cholerae* tends to be more numerous in the environment during warmer months.

There are a number of types of *V. cholerae*, and these produce very different symptoms. One type, *V. cholerae* 01, initially causes abdominal discomfort and mild diarrhea. As the illness progresses, the symptoms may include watery diarrhea, abdominal cramps, vomiting and dehydration. Death can occur. Susceptibility to cholera is enhanced in people who have had gastric surgery, take antacids or have type O blood.

**Slide 15**

Some controls for *Staphylococcus aureus* in seafood:
- Proper sanitation to prevent product contamination (product source, process facilities and personnel)
- Proper refrigeration to prevent growth
- Proper cooking
- Prevent cross-contamination after cooking
Another type of *V. cholerae*, non-01, causes diarrhea, abdominal cramps and fever. Nausea, vomiting and bloody diarrhea have also been reported. The severity of the symptoms is dependant, in part, upon the specific strain. In its most severe form, *V. cholerae* non-01 has resulted in septicemia (blood poisoning) in individuals with medical conditions that weaken their immune systems. The illness has been associated with consumption of raw oysters, but the bacterium has also been found in blue crabs.

*V. cholerae* can be prevented by cooking seafood thoroughly and by preventing cross-contamination once the seafood is cooked (Slide 16).

**Vibrio parahaemolyticus (Non-sporeformer)**

*Vibrio parahaemolyticus* is naturally occurring in estuaries and other coastal areas throughout most of the world. In most areas, *V. parahaemolyticus* is more numerous in the environment during the warmer months and, as a result, most outbreaks in the United States occur during the summer.

The most commonly experienced symptoms of *V. parahaemolyticus* illness include diarrhea, abdominal cramps, nausea, vomiting and headache. Fever and chills are less frequently reported. The illness has been associated with consuming contaminated crabs, oysters, shrimp, and lobster.

Hazards from *V. parahaemolyticus* can be controlled by thoroughly cooking seafood and preventing cross-contamination after cooking. Control of time/temperature abuse is also an important control measure (Slide 16).

**Vibrio vulnificus (Non-sporeformer)**

*Vibrio vulnificus* is a naturally occurring marine bacterium. *V. vulnificus* requires salt for survival and is commonly isolated at salinities of 7 parts per thousand (ppt) to 16 ppt. It is primarily found in the Gulf of Mexico, but it has also been isolated from the Atlantic and Pacific Oceans. The numbers of the bacterium in the environment are highest during the warmer months of April through October.

The most common symptoms include: skin lesions, septic shock, fever, chills and nausea. Abdominal pain, vomiting and diarrhea are less frequently reported. Death occurs in about 50 percent of the cases. A number of medical conditions make individuals more susceptible to the life-threatening effects of this bacterium, including liver disease, alcohol abuse, cancer, diabetes, chronic kidney disease, immunosuppressive drug or steroid usage, low stomach acidity, and AIDS. *V. vulnificus* sepsis has been associated with the consumption of certain molluscan shellstock.

Hazards from *V. vulnificus* can be controlled by thorough cooking of shellfish and by preventing cross-contamination once the seafood is cooked. The risk of *V. vulnificus* infection may also be reduced by refrigerating oysters soon after harvest during warm-weather months. Individuals in the “high risk” groups should not consume raw molluscan shellfish (Slide 16).
Viral Hazards

Like other microorganisms, viruses exist everywhere (Slide 17). They are very small particles that cannot be seen with a traditional microscope and cannot reproduce by themselves. Viruses exist in foods without growing, so they need no food, water or air to survive. They do not cause spoilage. Viruses cause illness by infection. They can infect living cells and reproduce inside the host cell. Viruses only grow once they enter a suitable host, and only some viruses consider humans a suitable host. Viruses can survive in human intestines, contaminated water, and frozen foods for months.

Viruses can be transmitted to infect consumers by contact with people, food or contaminated waters. They can be found in people who were previously infected but are no longer ill. Viruses can also be present in people who show no outward signs of illness (carriers). Transmission of viruses to foods is usually related to poor hygienic practices or harvest from non-approved, contaminated waters (e.g., illegal shellfish harvest). People who have viruses shed the particles when they defecate. Food handlers with viruses can transmit them to food if they do not wash their hands properly. Poor hygienic practices can also result in contamination of food with bacterial hazards as well as viral hazards.
The following are examples of viral hazards that can be found in seafood (Slide 18).

**Slide 18**

**Viruses:**

- Hepatitis A virus causes fever and abdominal discomfort, followed by jaundice.
- Norovirus group (formerly Norwalk Virus) causes nausea, vomiting, diarrhea, and abdominal pain (gastroenteritis). Headache and low-grade fever may also occur.

**Hepatitis A Virus**

Viruses survive better at low temperatures and are killed at high temperatures. As a result, most outbreaks of hepatitis occur during winter and early spring. Viruses can remain alive for long periods of time in seawater and have been shown to survive over one year in marine sediments.

Both raw and steamed clams, oysters, and mussels have been implicated in outbreaks of hepatitis A. Symptoms of hepatitis A include weakness, fever and abdominal pain. As the illness progresses, the individual usually becomes jaundiced. The severity of the illness ranges from very mild (young children often experience no symptoms) to severe, requiring hospitalization. The fatality rate is low, and deaths primarily occur among the elderly and individuals with underlying diseases.

Hazards from hepatitis A can be prevented by thoroughly cooking seafood and by preventing cross-contamination of cooked seafood. However hepatitis A appears to be more resistant to heat than other viruses. A laboratory study showed that hepatitis A viruses in infected oysters were inactivated after heating at 140°F for 19 minutes. Therefore, mollusks steamed only until the shells open (a common cooking practice) are not exposed to heat long enough to inactivate hepatitis A viruses (Slide 19).

**Norovirus**

Norovirus is considered a major cause of nonbacterial intestinal illness (gastroenteritis). Illness from Norovirus has been associated with eating raw and steamed clams, oysters and cockles. Norovirus causes nausea, vomiting, diarrhea, abdominal cramps, and occasionally fever.

Hazards from Norovirus can be prevented by properly cooking seafood and by preventing cross-contamination of cooked seafood. Additionally, a recent outbreak has demonstrated that controlling overboard discharge of untreated sewage from shellfish harvesting vessels would reduce the incidence of illness attributable to Norovirus (Slide 19).
Parasites are organisms that need a host to survive, by living on the host or within the host. Thousands of kinds of parasites exist worldwide; however, only about 20 percent can be found in food or water, and less than 100 are known to infect people through consumption of contaminated food or water. There are two types of parasites that can infect people through food or water: parasitic worms and protozoa. Parasitic worms include roundworms (nematodes), tapeworms (cestodes) and flukes (trematodes). These worms vary in size from barely visible to several feet in length. Protozoa are single-cell animals, and most cannot be seen without a microscope (Slide 20).

For most foodborne parasites, the food is part of their natural life cycle (e.g., nematode worms in fish and meat). They have the opportunity to infect humans when people eat them along with the food. The two factors most important to parasitic survival are a proper host (i.e., not all organisms can be infected by parasites) and a suitable environment (i.e., temperature, water, salinity, etc.).

Some parasites may be transmitted through food or water that is contaminated by fecal material shed by infected hosts (Slide 21). Consumer exposure to parasites depends on food selection, cultural habits and preparation methods. Most parasites do not harm humans but may be aesthetically unpleasant. Parasitic infections are normally associated with raw or undercooked foods because thorough cooking of foods eliminates all foodborne parasites. In specific instances, freezing can be used to destroy parasites in food.

The seafood parasite hazards have usually involved certain ready-to-eat products such as ceviche (pickled fish marinated with lime juice), lomi lomi (fish in a lemon base marinade), sashimi (raw fish), sushi (raw fish with rice), and cold smoked fish. The hazard concerns will vary according to different fish species, size and location of harvest. The following are examples of parasite hazards that have been associated with certain seafood (Slide 22).
Anisakis simplex

Anisakis simplex, commonly called herring worm, is a parasitic nematode or roundworm. Its final hosts are dolphins, porpoises and sperm whales. The larval (wormlike) stage in fish and squid is usually 0.7 to 1.4 inches in length and appears pinkish to whitish in color.

Anisakiasis, the human illness caused by Anisakis simplex, is associated with eating raw fish (sushi, sashimi, lomi lomi, ceviche, sunomono, Dutch green herring, marinated fish and cold-smoked fish) or undercooked fish.

Parasites in fish are considered a hazard only in fish that the processor knows or has reason to believe will be served raw or undercooked. In other products, parasites are considered filth but not hazardous. The FDA has established three freezing processes to kill parasites. Freezing and storing at -4°F (-20°C) or below for 7 days (total time), or freezing at -31°F (-35°C) or below for 15 hours, or freezing at -31°F (-35°C) or below until solid and storing at -4°F (-20°C) or below for 24 hours is sufficient to kill parasites. FDA's Food Code recommends these freezing conditions to retailers who provide fish intended for raw consumption. Note: these conditions may not be suitable for freezing particularly large fish (e.g. thicker than six inches) (Slide 23).

Pseudoterranova decipiens

Pseudoterranova decipiens, commonly called “codworm” or “sealworm,” is another parasitic nematode or roundworm. The usual final hosts of Pseudoterranova are gray seals, harbor seals, sea lions and walruses. The larval stage in fish are 0.2 to 2.2 inches in length, appearing yellowish, brownish or reddish in color.
These nematodes are related to *Anisakis simplex* and the disease associated with infections is also termed anisakiasis. These nematodes are also transmitted to humans through raw or undercooked fish. Control of *Pseudoterranova* is the same as for *Anisakis simplex* (Slide 23).

**Diphyllobothrium latum**

*Diphyllobothrium latum* is a cestode, or tapeworm, that parasitizes a variety of fish-eating mammals of the northern latitudes. A similar species is found in the southern latitudes and is associated with seal hosts. Cestodes have a structure that allows them to attach to the intestinal wall of their host and have segmented bodies. Cestode larvae found in fish range from less than one inch to several inches in length and are white or gray in color.

Diphyllobothrium tapeworms primarily infect freshwater fish. But salmon and related fish can also carry the parasites. Diphyllobothrium tapeworms are usually found unencysted and coiled in musculature or encysted in viscera. These tapeworms can mature and cause disease in humans. These cestodes are also transmitted to humans through raw or undercooked fish. Control of Diphyllobothrium is the same as for *Anisakis simplex* (Slide 23).

Some controls for *Anisakis simplex, P. decipiens* and *D. latum* parasites in seafood:
- Proper freezing
- Proper cooking

**Flukes (Flatworms)**

Flatworms is the common term used to refer to a large number of potential parasitic worms that appear as small, flat, and slender worms measuring about 20 mm long and 3-5 mm wide. They are also described according to the human organs they can infect, e.g., liver flukes, lung flukes, and intestinal flukes. Pain and inflammation is associated with the area of infestation, which can progress to more complications and death. Collectively, they cause the largest number of seafoodborne parasitic infections in the world, but occurrence is primarily within endemic areas of Asia including Korea, China, Taiwan, Vietnam and some Pacific Islands. These infections involve consumption of raw, fermented, undercooked or improperly pickled fish, crustaceans, and some mollusks usually taken from freshwater sources that involve the life-cycle of the parasite. They are largely associated with cultural preferences and recipes for freshwater products, e.g., congee (rice gruel with raw fish, wine soaked crab, and crab juices used for flavoring). Freezing, and proper heating or cooking can be effective controls to prevent infections.
Chemical Hazards

Chemical hazards can include a large variety of risks from natural conditions, intentional use and unintentional, or accidental contamination (Slide 24). Likewise, chemicals can be either beneficial or harmful depending on the types and amount used with seafood. Table 2 at the end of the chapter lists the types of materials that can be chemical hazards in foods.

### Slide 24

- **Chemical hazard in seafood due to natural conditions**
  - Biotoxins from natural sources (shellfish poisonings, ciguatera, tetrodotoxins)
  - Toxics due to product composition (gempylotoxin)
  - Toxins produced by bacterial growth on certain seafood (elevated histamine levels or scombrotoxins)
- **Chemical hazards added intentionally but improperly**
  - Food additives or processing aids
  - Aquaculture drugs
- **Chemical hazards due to unintentional or accidental contamination** (potentially toxic compounds or ingredients)
- **Allergens** — certain proteins that could present a risk to consumers who are not aware of their presence

**Natural conditions** involve seafood exposure to other biological conditions that can contaminate the seafood such that it could be potentially toxic when consumed either raw or cooked. Some intentionally added chemicals can be helpful in processing or for controlling bacteria growth if they are used in the proper, specified manner. Potential risks to consumers can increase when certain added chemicals are not controlled or the recommended treatment rates are exceeded. Regulatory limits are set for some of those contaminants. In contrast, some chemicals can be unintentionally added by accident or lack of knowledge such that they pose a definite seafood safety hazard.

**Allergens** are grouped as potential chemical hazards, but they are usually considered processing hazards if they are not declared in labels or other product information.

### Chemical Hazards Due to Natural Conditions

Marine biotoxins (natural toxins) represent a significant threat to human health when humans consume fish and fishery products that contain certain small amounts of these toxins (Slide 25). The marine biotoxins comprise many distinct compounds, all produced by species of naturally occurring marine algae. Algae are at the base of the marine food chain. Consequently, the biotoxins produced by some algae are collected and concentrated through the food chain (e.g., mollusks, crustaceans and finfish) and ultimately are consumed by humans.

There are several recognized marine shellfish biotoxins; e.g. paralytic shellfish poisoning (PSP), neurotoxic shellfish poisoning (NSP), diarrhetic shellfish poisoning (DSP), and amnesic shellfish poisoning (ASP). Molluscan shellfish
waters are classified by state shellfish-control agencies to reduce the risk that these toxins will be carried by shellfish in commerce or from recreational harvest. Processors must assure production of molluscan shellfish only from those waters that have been approved for harvest. The geographic extent or existence of such shellfish toxins in waters outside the US would also be subject to evidence of careful monitoring for approved harvest.

FDA has established action levels for all of the marine biotoxins except CFP. None of these toxins can be fully destroyed by normal cooking, freezing, salting, acidification or smoking processes. However, there is some evidence that PSP levels, and perhaps levels of other shellfish toxins, can be reduced to safe levels through commercial canning processes.

**Amnesic Shellfish Poisoning (ASP)**

ASP is primarily caused by contaminated molluscan shellfish, primarily from cold water regions of North America. The shellfish become contaminated with domoic acid produced by dense growths of certain algae in the genus *Pseudonitzschia*. It should be assumed that all filter-feeding mollusks are capable of accumulating domoic acid. However, the only shellfish implicated in cases of ASP have been mussels. ASP has also been identified as a problem in the viscera of dungeness, tanner, and red rock crabs, and anchovies along the western coast of the U.S. and Canada.

In the early stages of ASP, the individual usually experiences intestinal distress. Severe ASP can cause a facial grimace or chewing motion, short-term memory loss and difficulty breathing. Death can occur. Controls for amnesic shellfish toxins in seafood largely depend on harvest of approved products from approved waters (Slide 26).

**Diarrhetic Shellfish Poisoning (DSP)**

DSP is caused by contaminated molluscan shellfish. There has been no documented occurrence to date in the United States. However, instances have been documented in Japan, Southeast Asia, Scandinavia, western Europe, Chile, New Zealand, and eastern Canada. Filter-feeding mollusks can accumulate toxins even at algae concentrations below that necessary to discolor the water. Mussels, oysters, hard clams and soft-shell clams have been implicated in
cases of DSP. Contaminated scallops have caused cases of DSP in Japan, but the likelihood of scallops causing illness in this country is reduced because roe-on scallops are not typically consumed in the United States. A number of algae species in the genus *Dinophysis* and *Prorocentrum* have been associated with DSP. These algae are responsible for the production of a number of toxins (okadaic acid and its derivatives).

The symptoms of diarrhetic shellfish poisoning are diarrhea, nausea, vomiting, moderate to severe abdominal pain and cramps, and chills. No known fatalities have occurred, and total recovery is expected within three days with or without medical assistance. Controls for diarrhetic shellfish toxins in seafood largely depend on harvest of approved products from approved waters (Slide 26).

**Neurotoxic Shellfish Poisoning (NSP)**

*Gymnodinium breve* was first recognized as causing NSP in the mid 1960s. Blooms of this algae usually result in fish kills and can make shellfish toxic to humans. The blooms generally begin offshore and move inshore. *G. breve* produces three known toxins (brevetoxins).

NSP is caused by contaminated shellfish from the southeastern United States and New Zealand. Oysters and clams are the only shellfish associated with NSP illness. However, all filter-feeding mollusks are capable of accumulating neurotoxic shellfish toxins.

NSP resembles a mild case of ciguatera or PSP. Symptoms begin within three hours of consuming contaminated shellfish and include tingling of the face that spreads to other parts of the body, cold-to-hot sensation reversal, dilation of the pupils and a feeling of inebriation. Less commonly, victims may experience prolonged diarrhea, nausea, poor coordination, and burning pain of the rectum. Controls for neurotoxic shellfish toxins in seafood largely depend on harvest of approved products from approved waters (Slide 26).

**Paralytic Shellfish Poisoning (PSP)**

There are many species of toxic algae that cause paralytic shellfish poisoning. These include algae in the genus *Alexandrium*, *Pyrodinium* and *Gymnodinium*. PSP can be caused by a combination of any of 18 toxins (saxitoxins), depending on the species of algae, geographic area and type of shellfish involved.

PSP is caused by contaminated shellfish primarily from the U.S. Northeast and Northwest and imports from similar climates. All filter-feeding mollusks accumulate paralytic shellfish toxins. Mussels become highly toxic within a few hours to a few days of exposure to the organism, but also lose their toxin load rapidly. Clams and oysters generally do not become as toxic as mussels. They require more time to accumulate high levels of toxins and also require longer to cleanse themselves. Scallops can become extremely toxic, even during periods when blooms are not evident. However, scallops generally do not pose a PSP threat because the adductor muscle, the only part of the scallop traditionally consumed in the United States, does not accumulate toxin. PSP has recently been reported in the liver of Atlantic mackerel, American lobsters, and cold-water crabs such as dungeness, tanner, and red rock crab.
Symptoms of PSP initially involve numbness and a burning or tingling sensation of the lips and tongue that spreads to the face and fingertips. This leads to general lack of muscle coordination in the arms, legs and neck. A variety of other less commonly reported symptoms also exist. Severe cases of PSP have resulted in respiratory paralysis and death. Controls for paralytic shellfish toxins in seafood largely depend on harvest of approved products from approved waters (Slide 26).

**Slide 26**

Control for shellfish biotoxins in seafood:
- Only harvest approved shellfish products from approved waters.

**Ciguatera Fish Poisoning (CFP)**

One additional natural toxin of significant concern in the U.S. and about the world is ciguatera fish poisoning (CFP). Although it originates in a form of natural algae, the food route to humans is by toxin accumulation in certain fish harvested from certain areas. By eating toxic algae, certain species of tropical and subtropical fish can become toxic to humans. The algae species most often associated with CFP is *Gambierdiscus toxicus*, but others are occasionally involved. Toxic algae populations tend to fluctuate, influenced by the turbidity and nutrient content of the water. There are at least four known toxins that concentrate in the viscera, head or central nervous system of affected fish. Ciguatoxin is the principal toxin which can be in a variety of forms.

CFP has been carried to humans by contaminated finfish from the extreme southeastern United States, Hawaii, the tropics, and subtropics worldwide (between 35°N and 34°S latitude). In southern Florida, Bahamian and Caribbean regions, barracuda, amberjack, horseye jack, black jack, other large species of jack, king mackerel, certain large groupers and snappers have been associated with ciguatoxin. Many other species of large fish-eating fish may be suspect. In Hawaii and throughout the central Pacific, barracuda, amberjack and snapper are frequently ciguatoxic, and many other species, both large and small, may be suspect. Mackerel and barracuda from mid to northeastern Australian waters are frequently ciguatoxic.

The incidence of poisonous fish is sporadic. Not all fish of the same species and caught in the same area will necessarily be toxic. A study done in Hawaii indicated that if fish in one location are toxic, other fish in the vicinity are 60 percent likely to be toxic. Both plant-eating and fish-eating fish can become toxic. Plant-eating fish become toxic by eating the toxic algae itself. Fish-eating fish become toxic by consuming toxic plant-eating fish. Large fish are more likely to be poisonous than small fish because they consume greater amounts of the toxins.

Ciguatera causes diarrhea, abdominal pain, nausea, vomiting, abnormal or impaired skin sensations, vertigo, lack of muscle coordination, cold-to-hot
sensation reversal, muscular pain and itching. Some of the symptoms may recur for as long as six months. Death occasionally results.

Currently, the principal test method is a mouse bioassay that is not suitable for commercial use. There is no validated method suitable for shipboard or dockside testing of large catches of fish. However, some tests are being evaluated and may soon be available. In the meantime, the fishing industry must rely on local knowledge of safe harvest areas and avoid harvest from any officially designated areas or species (Slide 27).

**Slide 27**

Control for ciguatera in seafood:
- Do not process certain fish harvested from waters that have been designated as potentially ciguatoxic.

**Other Marine Toxins**

**Tetrodotoxin (puffer fish)**

Puffer fish, also called fugu or blowfish, contain the potent toxin, tetrodotoxin. It is unclear whether the fish itself produces the toxin, or like ciguatera, it is introduced to the fish by eating toxic algae. There are approximately 80 species of puffer fish that are known to contain tetrodotoxin in the Pacific, Atlantic and Indian Oceans. The domestic species of puffer, sometimes called sea squab, is much less poisonous than the Japanese species.

Symptoms of poisoning usually begin within 10 minutes of consuming puffer fish. The victim first experiences numbness and tingling of the mouth. This is followed by weakness, paralysis, decreased blood pressure, and quickened and weakened pulse. Death can occur within 30 minutes.

The primary control is avoidance of potential tetrodotoxic puffer fish (Slide 28). Puffer fish may not be imported into the United States except under strict certification requirements and specific authorization from FDA.

**Slide 28**

Control for tetrodotoxin in seafood:
- Do not process certain fish (puffer fish) that have been designated as potentially tetrodotoxic.

**Gempylotoxins**

The gempylids, escolars or pelagic mackerels are a small group of fish-eating oceanic fish. Important species in this group include: *Lepidocybium flavobrunneum* (escolar — California, Peru, Hawaiian Islands, Australia, South Africa, Cuba, Aru Islands, Madeira), and *Ruvettus pretiosus* (oilfish, castor oil fish, purgative fish — tropical Atlantic and Indo-Pacific Oceans).
Gempylids produce an oil that has a purgative effect. The diarrhea caused by eating the oil contained in the flesh and bones of gempylid fish develops rapidly and is pronounced but generally without pain or cramping. No other bad effects have been reported. There are not specific legal restrictions, but authorities advise caution that gempylid fish should not be imported or marketed in the United States (Slide 29).

**Control for gempylotoxin in seafood:**
- Do not process certain potentially gempylotoxic fish.

**Scombrototoxin (fish histamine poisoning)**

Scombroid poisoning, also known as histamine poisoning, is caused by eating fish of certain species that have undergone some degree of spoilage by certain types of naturally occurring bacteria. These bacteria produce an enzyme that reacts with natural components (amino acids) of the fish flesh to produce histamine and other biogenic amine compounds (putrescine and cadaverine). Fish that have been involved in scombroid poisonings include tuna, mahi-mahi, bluefish, sardines, amberjack and mackerel. The toxin is not eliminated by cooking or canning.

Scombroid toxicity is a common illness associated with certain seafood. Illnesses are commonly reported each year and they are usually self-limiting (less than 24-hour duration). Deaths are rarely reported. Symptoms of scombroid poisoning begin within 30 minutes up to four hours of eating contaminated fish. The most common symptoms include a metallic, sharp or peppery taste; nausea; vomiting; abdominal cramps; diarrhea; swelling and flushing of the face; headache; dizziness; heart palpitations; hives; rapid and weak pulse; thirst; and difficulty in swallowing.

The histamine-forming bacteria usually grow rapidly only at high temperatures. At 90°F (32.2°C), unsafe levels of histamine may appear within six hours. At 70°F (21°C) the toxic conditions may appear within 24 hours. Because wide variations occur between individual fish even under the same conditions, it is necessary to consistently remove heat rapidly from the freshly harvested fish and maintain a low temperature until the fish are prepared for consumer use. Particularly for large fish, special precautions and equipment are required for the rapid removal of heat. Sensory analysis is a screening method that can help to reduce the risk of accepting histamine-containing fish. Periodic increases in product temperature during storage can result in more histamine being formed, but histamine may form during high temperature short time abuse without the usual odors of decomposition. Chemical analysis for histamine is also possible. A detailed knowledge of the temperature history of the product provides the best control measure (Slide 30).
Intentionally Added Chemicals – Added Chemicals or Ingredients

Intentionally added chemicals include various compounds or ingredients that are approved or recognized for use with seafood products or processes, but they must be used in an appropriate or specified manner based on good manufacturing practices, regulatory limits and/or expert advice. Likewise, they should comply with established food grade standards or guidelines that assure safe composition and sources. Improper use or use of compounds from improper sources can result in seafoodborne hazards that cause intoxication, allergic-type reactions or food intolerances. Some intentionally added chemicals used with seafood include food additives for preservation or processing aids, nutritional additives, and color additives (Slide 31).

Food and color additives are used in many fish and fishery products, including some usage by fishermen and aquaculturists. Many additives are acceptable in such products when used in conformity with GMPs and established limits. Other additives are not permitted in fish or fishery products. Before using a food additive, the processor should become familiar with the applicable legal limitations for its use. The processor should be especially aware of food additives that are known to cause allergic-type reactions or are otherwise linked to adverse health consequences if not properly used. These reactions can be severe (e.g., anaphylactic shock induced by sulfites or yellow 5 and 6 can be fatal). The use of color additives that are permitted should be carefully controlled to ensure they remain within established limits. Correct listing of food and color additives on the product label is a legal requirement.

Certain food and color additives can cause an allergic-type reaction (food intolerance) in consumers. Examples of such food and color additives that are used on fish and fishery products include sulfiting agents and FD&C Yellow #5. Sulfiting agents are mostly used during on-board handling of shrimp and lobster to prevent the formation of “black spot.” They are sometimes used by cooked octopus processors as an antioxidant, to retain the red color of the octopus.
skin. FD&C Yellow #5 is used during in-plant processing. These food and color additives are permitted for use in foods—with certain restrictions—but their presence must be declared on the label. This label declaration is particularly important to sensitive individuals.

Certain other food and color additives are prohibited from use in food because of a determination by FDA that they present a potential risk to the public health. Examples of such food and color additives include safrole and FD&C Red #4.

These chemicals are intentionally added to food at some point during the food’s growth, processing or distribution. Intentionally added chemicals are safe when used at established safe levels but can be dangerous when those levels are exceeded. Controls to prevent seafood hazards due to intentional use include monitoring for proper use and testing for resulting residuals, plus labeling information to alert certain consumers with food intolerances (Slide 32).

Controls for intentionally added chemicals in seafood:
- Use proper type and amount of chemicals.
- Label product to inform consumers (e.g., sulfites)

### Intentionally Added Chemicals – Aquaculture Drugs

Animal drugs may be used in the raising of aquatic species to: 1) treat or prevent disease, 2) control parasites, 3) affect reproduction, and 4) tranquilize. Illegal residues of drugs may occur in aquaculture species because of the use of unapproved drugs, use of drugs not in accordance with the approved labeling directions, failure to follow approved withdrawal times, or use of general purpose chemicals not labeled or approved for drug use. There are only a few approved drugs for aquatic species. However, FDA approval is required before any animal drug is used to ensure that unsafe drug residues will not occur in edible tissue when animals are treated following approved label directions. The withdrawal period is the period from the last time of drug treatment until the residuals are reduced or eliminated in the edible portions. The withdrawal time is usually within a number of days, depending on the drug, dosage, and growth of the seafood. Producer quality-assurance programs provide information and guidance for proper use of approved compounds and record-keeping practices that can be referenced in processor HACCP plans. Processors may consider conducting on-site audits of the animal-drug controls used by their producers. If rapid screening tests are considered for use by the processor or producer to detect or monitor drug residues in aquatic species, they must be validated for their intended use. These tests should only be used as a part of a complete risk-reduction, quality-assurance program and not be used as the only monitoring tool. Presently, FDA has no data to indicate these tests will provide reliable, quantitative results for drug screening in farm-raised aquatic species (Slide 33).
Unintentional or Incidental Chemical Contamination

Chemicals can become part of a food without being intentionally added. These incidental chemicals might already be in a food ingredient when it is received. For example, certain seafood may contain small but legal residues of approved antibiotics. Packaging materials that are in direct contact with ingredients or the product can be a source of incidental chemicals, such as sanitizers or inks. Most incidental chemicals have no effect on food safety, and others are only a concern if they are present in excessive amounts. Incidental chemicals also include accidental additions of prohibited substances such as poisons or insecticides that may not be allowed at any level (Slide 34).

Unintentionally or Incidentally Added Chemicals:

- Agricultural chemicals (e.g., pesticides, herbicides, fungicides, antibiotics, growth hormones) can be acutely toxic if present in the food at high levels and may cause health risks with long-term exposure.
- Cleaning chemicals (e.g., sanitizers, chlorine, acids, caustics) can cause chemical burns if present in the food at high levels.
- Maintenance chemicals (e.g., lubricants, paints) that are not approved for food use and may be toxic
- Prohibited substances and toxic elements (e.g., lead, zinc, arsenic, mercury, cyanide)
- Polychlorinated Biphenyls (PCBs)

Fish are harvested from waters that are exposed to varying amounts of environmental contaminants. Industrial chemicals, pesticides, and many toxic elements may accumulate in fish at levels that can cause public health problems. Of greatest concern are fish harvested from freshwater, estuaries, and near shore waters rather than from the open ocean. Pesticides and herbicides used near aquaculture operations are also of concern. Federal tolerances or action levels are established for some of the most toxic and persistent contaminants. States often use these limits for deciding whether to close waters for harvesting. Processors should be aware of these closures and should not purchase fish that have been harvested in closed areas. Pesticides and herbicides that may be used near aquaculture operations are also potential problems. Producer quality-assurance programs provide useful information for avoiding potential contaminants from a variety of sources, beginning with proper site selection.
Controls are necessary to prevent potential hazardous chemical contaminations (Slide 35).

**Slide 35**

Some controls for unintentional or incidental chemical contamination of seafood:
- Proper use of cleaning and maintenance chemicals in the processing areas
- Proper location and monitoring of aquaculture farming operations relative to potential land runoffs and spraying of hazardous chemicals
- Do not harvest from polluted or non-approved waters
- Product screening relative to source

**Allergens**

A number of foods contain allergenic proteins that can pose a health risk to certain sensitive individuals. Foods that account for most of all food allergies include peanuts, soybeans, milk, eggs, fish, crustaceans, tree nuts, and wheat (Slide 36). If these foods are part of, or are directly added to your fishery product, you must ensure that the product is properly labeled. However, these controls are not designed to prevent the unintentional introduction of allergenic proteins from such foods into your fishery product because of cross-contact (e.g. use of common equipment, improper production scheduling, or improper use of rework material). Unintentional introduction of allergenic proteins must be controlled through a rigorous sanitation regime, either as part of a prerequisite program or as part of HACCP itself. The basic controls for allergens involve product declarations and monitoring to prevent cross-contamination among foods (Slide 37).

**Slide 36**

Most common food allergens:
- Milk
- Peanuts
- Soybeans
- Eggs
- Tree Nuts
- Wheat
- Fish
- Crustaceans

**Slide 37**

Control for potential allergens in seafood:
- Product labeling to inform consumers
Physical Hazards

Physical hazards include any potentially harmful extraneous matter not normally found in food (Slide 38). When a consumer mistakenly eats the foreign material or object, it is likely to cause choking, injury or other adverse health effects. Physical hazards are the most commonly reported consumer complaints because the injury occurs immediately or soon after eating, and the source of the hazard is often easy to identify. Table 3 at the end of the chapter lists the types of materials that can be physical hazards in foods.

**Slide 38**

Physical Hazard:
Any extraneous matter not normally found in food that could cause physical injury.

**Example:**

The following are examples of materials that may be physical hazards:

<table>
<thead>
<tr>
<th>Material</th>
<th>Why a hazard?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glass</td>
<td>Cuts, bleeding; may require surgery to find or remove</td>
</tr>
<tr>
<td>Metal</td>
<td>Cuts, broken teeth; may require surgery to remove</td>
</tr>
</tbody>
</table>

**Glass Inclusion Hazards**

Glass fragments can cause injury to the consumer. FDA's Health Hazard Evaluation Board has supported regulatory action against products with glass fragments of 0.3 inches (7 mm) to 1.0 inches (25 mm) in length. Glass inclusion can occur whenever processing involves the use of glass containers. Normal handling and packaging methods, especially mechanized methods, can result in breakage. Most products packed in glass containers are intended as a ready-to-eat commodity. Glass fragments originating from other sources must be addressed, where applicable, in a prerequisite sanitation program (Slide 39).

**Note**

Bone and shell fragments are typically considered an intrinsic part of the seafood products (fish fillets and shellfish meat) and not a contaminant. Hazard analysis considers them as quality problems and not a significant safety hazard.

**Slide 39**

Control for potential glass inclusion in seafood:
- Examination of glass containers for breakage
Metal Hazards

Metal-to-metal contact—especially in mechanical cutting and blending operations, and with equipment that has parts that can break or fall off, such as wire-mesh belts—can introduce metal fragments into products. FDA's Health Hazard Evaluation Board has supported regulatory action against products with metal fragments of 0.3 inches (7 mm) to 1.0 inches (25 mm) in length. Such fragments serve as a hazard to the consumer. This hazard can be controlled by subjecting the product to metal detection devices or by regular inspection of at-risk equipment for signs of damage. Controls to prevent potential inclusion of metal fragments include monitoring of the equipment and detection in products (Slide 40).

Slide 40

Controls for potential metal inclusion in seafood:
• Monitoring equipment for wear and breakage
• Screening products with metal detectors

Table 1

Types of Biological Hazards
1) Bacteria
   a) Sporeformers
      – Clostridium botulinum
      – Clostridium perfringens
      – Bacillus cereus
   b) Nonsporeformers
      – Campylobacter spp.
      – Listeria monocytogenes
      – Salmonella spp. (e.g., S. typhimurium, S. enteritidis)
      – Shigella spp. (e.g., S. dysenteriae)
      – Staphylococcus aureus
      – Vibrio spp. (e.g., V. cholerae, V. parahaemolyticus, V. vulnificus)
      – Yersinia enterocolitica

2) Viruses
   – Hepatitis A and E
   – Norovirus group

3) Parasites
   – Anasakis simplex
   – Ascaris lumbricoides
   – Diphyllobothrium latum
   – Pseudoterranova dicepiens
Types of Chemical Hazards
1) Naturally Occurring Chemicals
   – Scombrotoksin (histamine)
   – Ciguatoxin
   – Gempylotoxins
   – Shellfish Biotoxins
      – Paralytic shellfish poisoning (PSP)
      – Diarrheic shellfish poisoning (DSP)
      – Neurotoxic shellfish poisoning (NSP)
      – Amnesic shellfish poisoning (ASP)/Domoic acid
   – Food Allergens
2) Intentionally Added Chemicals
   – Food additives
   – Preservatives (e.g., nitrite and sulfiting agents)
   – Nutritional additives (e.g., niacin)
   – Color additives
3) Unintentionally or Incidentally Added Chemicals
   – Agricultural chemicals (e.g., pesticides, fungicides, herbicides, fertilizers, antibiotics and growth hormones)
   – Prohibited substances (Code of Federal Regulations, chapter 21, section 189)
   – Toxic elements and compounds (e.g., lead, zinc, arsenic, mercury and cyanide)
   – Polychlorinated biphenyls (PCBs)
   – Plant chemicals (e.g., lubricants, cleaning compounds, sanitizers and paints)

Table 3

Types of Physical Hazards
1) Glass Inclusion (bottles, jars, other containers)
2) Metal Inclusion (machinery, wire, hooks, staples)
Chapter 5

Principle 1: Hazard Analysis

The first step in developing a HACCP plan is to identify all of the significant food safety hazards that are associated with the seafood product(s) and process(es), as well as the control measures. This procedure is called hazard analysis, the first principle of HACCP (Slide 1).

In this chapter you will learn how to:
• Conduct a hazard analysis
• Identify significant hazards
• Identify control measures

A hazard is defined as any biological, chemical or physical agent that is reasonably likely to cause illness or injury in the absence of control(s). The term hazard, when used in the context of HACCP, is limited to food safety concerns that could cause consumer illness or injury (Slide 2).

Definition: A hazard is any biological, chemical or physical agent that is reasonably likely to cause illness or injury in the absence of control(s).
Chapter 5

Not all potential hazards are significant. A significant hazard is one that is reasonably likely to occur and presents a health risk to the consumer if it is not controlled (Slide 3). Control measures must be identified for all significant hazards.

**Slide 3**

The hazard analysis is conducted to identify:
- All potential food safety hazards,
- Which of these hazards are significant, and
- Measures to control the significant hazards.

**How to Conduct a Hazard Analysis**

There is a sequence of steps that need to be completed when conducting a hazard analysis (Slide 4). Each step is an important part of the procedure. These steps will be discussed individually using a worksheet to document the results or conclusions.

**Slide 4**

There are five steps in a hazard analysis:
1) List process steps
2) Identify potential food safety hazards
3) Determine if the hazard is significant
4) Justify the decision
5) Identify control measure(s)

A standardized hazard analysis worksheet (Slide 5) is designed to ensure that all steps in the hazard analysis process are completed. A written hazard analysis is important because it is the best way to determine if there are significant food safety hazards that need to be controlled.

The worksheet is used to:

- List each of the process steps from the process flow chart (Column 1).
- List all potential species-related and process-related hazards that are identified during the hazard identification step (Column 2).
- Record the result of the hazard evaluation. A “Yes” or “No” answer to the question: “Is the potential food safety hazard significant?” is entered in this column (Column 3).
- Explain why the hazard is significant or not (Column 4).
- List control measures for those hazards that have been identified as significant and need to be controlled at a specific operational step (Column 5).

**Note**

Blank hazard analysis worksheets are in Appendix 2.
Set Up the Hazard Analysis Worksheet

Set up the hazard analysis worksheet by entering the firm’s name and address. Then enter the information that was gathered during the preliminary steps, including the product description, the method of storage and distribution, and the intended use and consumer. A separate worksheet may be needed for each product type. Grouping of product types may be done so long as the hazards and controls are the same.

A process flow chart was developed (Chapter 3) as part of the preliminary steps. List each of these process steps in Column 1 of the hazard analysis worksheet (Slide 6).

Slide 6

Step 1:
Enter the processing steps from the process flow chart.

List all of the potential food safety hazards related to each type of fish or fishery product and the process in column 2 of the hazard analysis worksheet. All of the potential hazards should be listed at each processing step (Slide 7).
One approach to identify potential seafood safety hazards is to use the FDA Fish and Fisheries Products Hazards and Controls Guidance (Hazards Guide). It lists all of the potential seafood safety hazards that are likely to be associated with specific species of fish and shellfish and specific types of finished products or processing operations. The Hazards Guide is based on the best currently available scientific information (Slide 8).

### Slide 7

**Step 2:**
List potential food safety hazards.

### Using the Hazards Guide to Identify Potential Seafood Safety Hazards

The Hazards Guide provides tables of information in Chapter 3 that can be used to identify potential species- and process-related hazards.

#### Species-related Hazards

The Hazards Guide’s “potential vertebrate species-related hazards” table contains a list of all species of vertebrate fish (fish with backbones) in alphabetical order (Slide 9). The first column lists the “market names” for each type of fish. FDA’s “The Seafood List” contains the acceptable market names for all fish and shellfish species in commerce. These market names can be matched with common and regional names.

The scientific or Latin name for each type of fish is listed in the second column. The scientific name consists of two Latin words in italics. The first word is capitalized and denotes the “Genus” name and the second designates the “species.” The scientific name is universally recognized throughout the world. This name may be needed to properly identify the species of fish being considered to ensure that the correct food safety hazards are identified. In some cases the fish will be grouped by genus only in the Hazards Guide. Slide 7 lists the scientific name for mahi-mahi as “Coryphaena spp.” This means that all species (spp.) in the Genus Coryphaena are covered by this entry.

The remaining columns in the Hazards Guide list the potential hazards that are known to be associated with vertebrate fish. These hazards are referred to as “species-related hazards.” If the potential hazard is reasonably likely to occur in a certain species of fish, there is a checkmark in the column for that particular hazard. This means that this hazard should be listed as a “potential” hazard in column 2 of the hazard analysis worksheet. Under each hazard at the top of

---

**Web Link**

**Seafood List**
http://www.fda.gov/Food/GuidanceComplianceRegulatoryInformation/GuidanceDocuments/Seafood/ucm113260.htm
Principle 1: Hazard Analysis

this chart, "CHP" refers to the chapter in the Hazards Guide that describes that hazard.

The Hazards Guide's "potential invertebrate species-related hazards" table (not shown in this manual) lists the same information for species of invertebrate fish (fish without backbones) in alphabetical order. All edible species of bivalve shellfish, mollusks, and crustaceans are included in this table.

Slide 9

Identifying Potential Species-Related Hazards

<table>
<thead>
<tr>
<th>Market Names</th>
<th>Latin Names</th>
<th>Hazards</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Parasites</td>
<td>Natural Toxins</td>
<td>Histamine</td>
<td>Environmental Chemicals</td>
<td>Aquaculture Drugs</td>
</tr>
<tr>
<td>Mahi-mahi</td>
<td><em>Coryphaena</em> spp.</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mahi-mahi, aquacultured</td>
<td><em>Coryphaena</em> spp.</td>
<td>✓ ✓ ✓</td>
<td></td>
<td>✓ ✓ ✓</td>
<td>✓ ✓ ✓</td>
<td></td>
</tr>
<tr>
<td>Marlin</td>
<td><em>Makaira</em> spp.</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Tetrapturus</em> spp.</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Process-related Hazards

The Hazards Guide's "potential process-related hazards" table lists the potential hazards that are associated with all finished product forms and package types (Slide 10). This table is also formatted in columns. The first column lists the "finished product food types." The second column lists the "package type," which is divided into three categories: a) “reduced-oxygen packaged” (ROP) – products that are packaged in a reduced-oxygen environment, b) “other than reduced-oxygen packaged” – products that are not, and c) “all” – products that can be either reduced-oxygen packaged or not. The remaining columns in the table list all of the potential process-related food safety hazards that might occur in seafood products or processing operations. When selecting a finished food and package type it is important to review all of the entries in this table and look for the best fit for the product being considered.
The Hazards Guide’s “potential process-related hazards” table was used to identify potential species-related hazards for the fresh mahi-mahi fillets received by XYZ Seafood Company. XYZ Seafood Company is receiving wild-caught mahi-mahi fillets from another processor. The hazard of histamine is checked and should be listed in Column 2 of the hazard analysis worksheet. This table is used in the same way to determine potential food safety hazards for any species of fish.

To complete the hazard identification, find the process-related hazards for this product from the Hazards Guide’s “Potential Process-Related Hazards” table (Slide 10). Begin by identifying the finished product food description that best fits this product. For fresh mahi-mahi fillets, there is an entry for: *Raw fish other than oysters, clams, and mussels (finfish and non-finfish)*. This entry is the best match for the fresh mahi-mahi fillets.

The next step is to determine whether or not this product is received, stored or placed in a reduced-oxygen package while in this firm’s control. If no reduced-oxygen packaging is used, select the entry for “*Raw fish other than oysters, clams, and mussels + other than reduced-oxygen packaged*.” Hazards Guide Table #3-4 shows that there are three potential food safety hazards for this product form:

- Pathogen growth – temperature abuse (biological hazard)
- Allergens and/or additives (chemical hazards)
- Metal inclusion (physical hazard).

The pathogen growth-temperature abuse hazard has a superscript next to the checkmark that refers to a footnote at the bottom of the table. This footnote states that this hazard only applies if the product is intended to be consumed without cooking. To decide whether or not to include this hazard, consider who this product is being sold to and how it will be consumed. If the product is being sold to retail stores and/or restaurants for consumption by the general public, most processors can assume that the fillets will be cooked before they are consumed and do not need to list this hazard in the hazard analysis. However, if the product was not intended to be cooked by the consumer (for example, sushi, sashimi, a marinated or partially cooked product) this food safety hazard must be included in the hazard analysis.

Based on the XYZ Seafood Company process description on page 37, a hazard analysis worksheet (Slide 11) has been developed by entering:

- Company and product information at the top of the worksheet. This information was gathered in the preliminary steps (Chapter 3).
- The first step from the process flow chart (Receiving Fresh Fillets) was entered in column 1.
- The species-related hazard, histamine, found in Hazards Guide Table #3-2 for mahi-mahi was entered in column 2.
- The process-related hazards, food allergens and metal inclusion, found in Hazards Guide Table #3-4, for raw fish other than vacuum packaged were entered in column 2.

Note

If the product was vacuum-packed or packed in a reduced-oxygen package, you would select the “Reduced-oxygen packaged” entry in the hazards guide, which also identifies the hazard of “*C. botulinum* growth.”
### Identify potential process-related hazards

#### Hazards Guide’s Potential Process-Related Hazards Table

<table>
<thead>
<tr>
<th>Finished Product Food¹</th>
<th>Package Type</th>
<th>Hazards</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pathogenic bacteria growth – temperature abuse</td>
<td></td>
</tr>
<tr>
<td></td>
<td>C. botulinum toxin</td>
<td></td>
</tr>
<tr>
<td></td>
<td>S. aureus toxin – drying</td>
<td></td>
</tr>
<tr>
<td></td>
<td>S. aureus toxin – batter</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pathogens survival through cooking or pasteurization</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pathogens survival designed to retain raw product characteristics</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pathogenic bacteria contamination after pasteurization</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Allergens/Additives</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Glass inclusion</td>
<td></td>
</tr>
</tbody>
</table>

| Raw fish other than oysters, clams, and mussels (finfish and non-finfish) | Reduced-oxygen packaged (e.g. mechanical, vacuum, steam flush, hot fill, MAP², CAP², hermetically sealed or packed in oil) | ✓ √ | √ |
| Raw fish other than oysters, clams, and mussels (finfish and non-finfish) | Other than reduced-oxygen packaged | ✓ √ | √ |
| Partially cooked or uncooked prepared foods | Reduced-oxygen packaged (e.g. mechanical, vacuum, steam flush, hot fill, MAP², CAP², hermetically sealed or packed in oil) | ✓ √ | ✓ ✓ ✓ |
| Partially cooked or uncooked prepared foods | Other than reduced-oxygen packaged | ✓ | ✓ ✓ ✓ |

¹You may need to include potential hazards from more than one finished product food category if your product fits more than one description.
²MAP = Modified atmosphere packaging; CAP = Controlled atmosphere packaging.
³This hazard only applies if you have knowledge or have reason to know that the fish will be consumed without a process sufficient to kill pathogens, or if you represent, label, or intend for the product to be so consumed.

### Hazards Not Identified in the Hazards Guide

In addition to using the Hazards Guide to identify hazards, it is recommended that firms also perform an on-site assessment of the activities and conditions at each operational step. Firms may note unique product sources, conditions, activities, or new research information that is not identified in the Hazards Guide tables and could negatively impact product safety.

There may be situations where a firm identifies a hazard that is not listed in the Hazards Guide. For example, environmental chemical contaminants are not identified for many species. However, if a processor knows that fish they received are caught in an area that experienced a recent accidental spill of one or more chemical contaminants then this hazard should be considered in the hazard analysis.
Sanitation-related Hazards

The HACCP team may identify hazards caused by conditions or activities in the plant such as employee practices. Rather than controlling these sanitation-related hazards in a HACCP plan, it is recommended that the sources of contamination be controlled in a separate, but equally important, Sanitation Control Procedures (SCPs). A Sanitation Control Procedures (Chapter 2) applies to all parts of the operation and is designed to prevent the many different sources of contamination from the plant environment. Therefore, when evaluating hazards associated with contamination from the plant environment, the HACCP team can determine that it is not significant because it is being monitored and controlled by the SCP.

Hazard Evaluation and Justification

After the hazard identification (step 2) is completed, the HACCP team must evaluate the hazards (Slide 12). Each potential hazard will be evaluated to determine if it is significant at each process step. The hazard evaluation, or risk assessment, is designed to determine which hazards are reasonably likely to occur and need to be controlled.
Principle 1: Hazard Analysis

Steps 3 and 4: Hazard Evaluation and Justification – Determine which hazards are significant and explain why.

HACCP focuses solely on food safety hazards that are reasonably likely to occur and are likely to result in an unacceptable health risk to consumers if they are not controlled. The hazard evaluation is designed to determine which hazards are relevant (Slide 13).

To determine if a hazard is significant, consider two questions:
1) Is the hazard reasonably likely to occur in the finished product in the absence of control?
2) Is the hazard likely to cause consumer illness?

Some processors will have the expertise necessary to complete the hazard evaluation, while others may need to seek outside assistance to complete this step. The HACCP team should use the Hazards Guide, experience, and other tools (e.g. test results, studies, FDA Alerts, recalls, etc.) that are available to help them determine whether or not a hazard is significant. There may be differences of opinion, even among experts, regarding whether or not a hazard is significant in a given situation.

Example: Fresh Mahi-mahi/XYZ Seafood Co.

Hazard Evaluation and Justification (Columns 1 – 4)

The potential hazards identified for XYZ Seafood Company (Slide 14) include:

- one species-related hazard: histamine, and
- two process-related hazards: allergens/additives, and metal inclusion.

Example – Fresh Mahi-mahi
Which Hazards are Significant at the first process step, Receiving?
- Histamine (Yes or No?)
- Allergens/Additives (Yes or No?)
- Metal Inclusion (Yes or No?)
Each of these potential hazards must be evaluated for significance at each processing step listed on the hazard analysis worksheet. The hazard evaluation will determine whether or not this hazard is likely to result in an unacceptable health risk to consumers if it is not properly controlled at each processing step.

**Process Step:** Receiving Fresh Fillets  
**Hazard:** Histamine

The Hazards Guide identified histamine as a species-related hazard in mahi-mahi that is reasonably likely to occur. To determine if this potential hazard could be introduced, enhanced (made worse), or eliminated at this process step, use the information in Chapter 7 of the Hazards Guide.

This chapter of the Hazards Guide states that histamine can form in certain species of fish, including mahi-mahi, when they are subject to time and temperature abuse. If abuse occurs at any time, from the time the fish is caught until the consumer eats it, this hazard could develop and cause illness. The Hazards Guide also states that histamine cannot be removed or eliminated once it has developed in the fish. This hazard must be **prevented** by making sure that these fish are not exposed to temperatures above 40°F for an extended period of time.

To determine if histamine is a **significant** hazard at the receiving step, answer two questions:

1) **Is the histamine hazard reasonably likely to occur in the absence of control?** The answer is “yes” because the Hazards Guide indicates that histamine is likely to develop in species like mahi-mahi if they are temperature-abused.

2) **If not properly controlled (in the process) is it likely to result in an unacceptable health risk to consumers?** Again, the answer is “yes” because at receiving, controls are needed to ensure that temperature abuse has not occurred during transit.

**Conclusion:** Since the answer to both of the hazard evaluation questions is yes, the hazard of histamine is significant at the receiving step.

**Process Step:** Receiving Fresh Fillets  
**Hazard:** Food Allergens/Additives

To determine if food allergen is a **significant** hazard in the process, answer two questions:

1) **Is the food allergen hazard reasonably likely to occur in the absence of control?** The answer is “yes” because the Hazards Guide (Chapter 19) indicates that finfish is one of the eight major food allergens.

2) **If not properly controlled (in the process) is it likely to result in an unacceptable health risk to consumers?** Again, the answer is “yes” because finfish is one of the eight major food allergens and can cause consumer illness.
Conclusion: Since the answer to both of the hazard evaluation questions is yes, the hazard of food allergens is significant at the receiving step.

Process Step: Receiving Fresh Fillets  
Hazard: Metal Inclusion

To determine if metal inclusion is a significant hazard at the receiving step, answer two questions:

1) Is the metal inclusion hazard reasonably likely to occur in the absence of control? The answer is “no” because it is not reasonably likely to occur for this process at any step because no metal is used so there is no chance for metal inclusion (Hazards Guide Chapter 20).
2) If not properly controlled (in the process) is it likely to result in an unacceptable health risk to consumers? Again, the answer is “no” because this hazard is not likely to occur.

Conclusion: Since the answer to both of the hazard evaluation questions is no, the hazard of metal inclusion is not significant at the receiving step.

Hazard Analysis Worksheet

The XYZ Seafood Company continued filling out the hazard analysis worksheet (Slide 15) by entering:

• Yes, in column 3 indicating that the species-related hazard of histamine is significant at the receiving step. In column 4 this decision is justified by stating that time/temperature abuse could occur during transit.
• Yes, in column 3 indicating that the process-related hazard of food allergens is significant at the receiving step. In column 4 the decision is justified by stating that fish is one of the eight major food allergens.
• No, in column 3 indicating that the process-related hazard of metal inclusion is not significant at the receiving step. In column 4 the decision is justified because it is not reasonably likely to occur.
Identifying Control Measures (column 5)

The final step in the hazard analysis process is to determine the appropriate control measures you intend to use to prevent, eliminate or reduce to an acceptable level each of the significant hazards identified in your hazard evaluation (Slide 16). This is important because if the hazard is truly significant there must be an appropriate control measure or your product would not be safe (Slide 17).

Control measures are actions and activities that can be used to prevent, eliminate, or reduce a food safety hazard to an acceptable level. In practice, control measures could include a wide array of activities that will be effective for a specific hazard.
Different control measures for biological hazards such as pathogenic bacteria, viruses and parasites may be used to control a hazard (Slide 18).

Control measures for pathogenic bacteria could include:

1) Time/temperature controls: proper control of refrigeration and storage time to minimize or prevent bacterial pathogen growth.
2) Heating and cooking processes to eliminate (kill) bacterial pathogens.
3) Freezing to prevent bacterial pathogen growth.
4) Fermentation and/or pH controls to ensure that foods are acidic enough to prevent bacterial pathogen growth.
5) Addition of salt or other preservatives to prevent bacterial pathogen growth.
6) Drying to ensure that enough water has been removed from the food to prevent bacterial pathogen growth.
7) Source control or buying raw material from acceptable sources to reduce the risk of bacterial pathogens to an acceptable level.
Control measures for pathogenic viruses could include:

1) Cooking methods designed to eliminate (destroy) viruses.
2) Source control or buying raw material from acceptable sources to reduce the risk of viruses to an acceptable level.

Control measures for parasites could include:

1) Cooking at the proper temperature for the proper amount of time to eliminate (kill) parasites.
2) Freezing at the proper temperature for the proper amount of time to eliminate (kill) parasites.

Different control measures for chemical and physical hazards may be used to control a hazard (Slide 19).

**Slide 19**

Control Measures for Chemical and Physical Hazards

**Chemical Hazards** *(Natural toxins, pesticides, drug residues, unapproved food and color additives, histamine)*

1) Source controls
2) Time/temperature controls
3) Production controls
4) Labeling controls

**Physical Hazards** *(Metal, glass, etc.)*

1) Source controls
2) Production controls

Control measures for chemical hazards, such as natural toxins, pesticides, drug residues, unapproved food and color additives, and histamine, could include:

1) Source controls to reduce the risk that fishery products were harvested from areas where chemical hazards like environmental pollutants or natural toxins like ciguatera are present at levels that could cause consumer illness or harm.
2) Time/temperature controls for chemical hazards such as histamine that prevent the hazard from forming in certain fish species.
3) Production controls to ensure that the proper amount of any food additives is used.
4) Labeling controls to ensure that consumers are aware that any known allergens like sulfites are present in the product.

Control measures for physical hazards, such as metal and glass, could include:

1) Source controls that reduce the risk that products supplied by vendors will contain any physical hazards like metal or glass.
2) Production controls such as visual inspections of the equipment or container, using magnets, metal detectors, sifter screens or other devices to prevent any finished products that contain metal or other physical hazards from entering the marketplace.
The XYZ Seafood Company continued the hazard analysis worksheet by adding control measures (column 5) for the identified hazards (Slide 20):

Slide 20

XYZ Seafood Company – Fresh Mahi-mahi Fillets

<table>
<thead>
<tr>
<th>Processing Step</th>
<th>Hazard</th>
<th>Significant (Yes or No)</th>
<th>Justify the decision</th>
<th>Control measure(s)</th>
<th>Critical Control Point?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receiving Fresh Fillets</td>
<td>Histamine</td>
<td>Yes</td>
<td>Time/temp. abuse during transit could cause histamine to form in the fish</td>
<td>Mahi-mahi fillets are shipped in containers buried in ice (proper icing)</td>
<td>No</td>
</tr>
<tr>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step (proper labeling)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metal Inclusion</td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- The mahi-mahi fillets will be received in containers buried in ice, to prevent time/temperature abuse during transit.
- The mahi-mahi fillets will be properly labeled at a later processing step, to control the food allergen hazard.

Summary

The hazard analysis is important because the decisions that are made will determine what is included in the HACCP plan. Tools, like the FDA's Hazards Guide, along with some thought and discussion, will ensure that the hazard analysis is completed successfully.
Chapter 5

The hazard analysis worksheet (see Appendix 2) can be used to document the decisions that are made. This documentation is not required by regulations, but it is strongly recommended for future reference and justification for the selection of hazards that are reasonably likely to occur and their controls.

Different Approaches and Teaching Strategies for Hazard Analysis

There are different approaches that can be used to conduct or teach hazard analysis (Slide 21). In the XYZ Seafood Company example presented in this chapter, the approach that is used relies on the Hazards Guide to determine the potential food safety hazards (inclusive method). All of these potential food safety hazards are then listed and evaluated at each processing step. Although it may be obvious that some of these hazards do not apply to some processing steps, this approach makes it less likely the hazards will be missed.

Slide 21

Different approaches in conducting a hazard analysis
- Traditional Method
- Inclusive Method

There are other approaches that can be used to conduct a hazard analysis. An alternative approach (traditional method) is to compile a list of all potential food safety hazards using the Hazards Guide and HACCP team experience. The team would use this list and decide which hazards are relevant to each processing step and list them only at that step on the hazard analysis worksheet. This approach will result in fewer hazards being listed at each processing step.

Either approach can be used to successfully complete a hazard analysis.

An example of the alternative approach to hazard analysis is provided for XYZ Seafood Company in Appendix 4 of this training manual.

Example: Fresh Mahi-mahi/XYZ Seafood Co.

Complete Hazard Analysis Worksheet

The hazard analysis worksheet summarizes the results of the completed hazard analysis for XYZ Seafood Company (Slide 22).
XYZ Seafood Company – Fresh Mahi-mahi Fillets

Hazard Analysis Worksheet

<table>
<thead>
<tr>
<th>Processing Step</th>
<th>(1)</th>
<th>(2)</th>
<th>(3)</th>
<th>(4)</th>
<th>(5)</th>
<th>(6)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process.</td>
<td>Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No)</td>
<td>Justify the decision that you made in column 3</td>
<td>What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard?</td>
<td>Is this step a Critical Control Point? (Yes or No)</td>
</tr>
<tr>
<td>Receiving Fresh Fillets</td>
<td></td>
<td>Histamine</td>
<td>Yes</td>
<td>Time/temp. abuse during transit could cause histamine to form in the fish</td>
<td>Mahi-mahi fillets are shipped in containers buried in ice (proper icing)</td>
<td></td>
</tr>
<tr>
<td>Food Allergens</td>
<td></td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step (proper labelling)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metal Inclusion</td>
<td></td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Refrigerated Storage</td>
<td></td>
<td>Histamine</td>
<td>Yes</td>
<td>Time/temperature abuse during storage could cause histamine to form in the fish</td>
<td>Mahi-mahi fillets are buried in ice &amp; stored in a refrigerated cooler (proper icing)</td>
<td></td>
</tr>
<tr>
<td>Food Allergens</td>
<td></td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step (proper labelling)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metal Inclusion</td>
<td></td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processing Step</td>
<td>(2) List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process.</td>
<td>(3) Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No)</td>
<td>(4) Justify the decision that you made in column 3</td>
<td>(5) What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard?</td>
<td>(6) Is this step a Critical Control Point? (Yes or No)</td>
<td></td>
</tr>
<tr>
<td>-------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Trim</td>
<td>Histamine</td>
<td>No</td>
<td>Not likely to occur, time at this and weigh/pack/label step is 30 minutes or less</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step (proper labelling)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Fillet knives are not likely to chip and contaminate product with metal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weigh/Pack/Label</td>
<td>Histamine</td>
<td>No</td>
<td>Not likely to occur, time at this and weigh/pack/label step is 30 minutes or less</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets are labeled with market name at this step (proper labelling)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Finished Product</td>
<td>Histamine</td>
<td>Yes</td>
<td>Time/temperature abuse could occur during storage</td>
<td>Mahi-mahi fillets are buried in ice &amp; stored in a refrigerated cooler (proper icing)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Refrigerated Storage</td>
<td>Food Allergens</td>
<td>No</td>
<td>Fillets were labeled with market name at Weigh/Pack/Label step</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Principle 2: Determine Critical Control Points

Introduction

This chapter will cover the second principle of HACCP – Critical Control Point (CCP) determination. For each significant hazard that was identified during the hazard analysis, there are one or more points or steps in the process where the hazard can be controlled. These points or steps are called Critical Control Points (Slide 1).

Slide 1

In this chapter you will learn:
- The definition of a Critical Control Point (CCP).
- The relationship between significant hazards, control measures, and CCPs.
- How CCPs may be different for different products and processes.
- Tools to help you determine which steps are CCPs.
- Examples of CCPs for various food safety hazards.

A CCP should be a specific point in the process flow where application of a control measure effectively prevents, eliminates or reduces the hazard to an acceptable level (Slide 2).

Slide 2

Definition: A Critical Control Point is a step at which control can be applied to prevent, eliminate a food safety hazard, or reduce it to an acceptable level.
Critical Control Point Placement

If there are no control measures that can be applied at a particular process step, that step cannot be the CCP. In some cases control measures should be applied at a particular step, but that step may not be the best place to control the hazard. In that case, a processing step that occurs later in the process flow may be the best place to control that hazard (Slide 3).

Slide 3

CCP placement must be at the processing step or steps that adequately control the significant hazard.

For example, when producing a cooked product, the hazard of pathogen growth would first be identified at the receiving step. This hazard could be controlled at any process step where the hazard can be adequately controlled. However, the best place to control this hazard would be the cooking step, where the hazard is eliminated. The cook step would be the CCP for this hazard.

There may be different control options for a single hazard. For example, a metal hazard can be controlled at different processing steps such as:

- At receiving, ingredients are sourced that are free of metal fragments,
- At a screening step, screens are used to remove any metal fragments,
- At a metal detection step, a detector is used to find any finished products that are contaminated with metal fragments.

Only one of these processing steps would likely be the best CCP to control this hazard.

Examples of Critical Control Points

A CCP is a step where a hazard can be prevented, eliminated or reduced to an acceptable level.

Examples of CCPs where a hazard can be prevented are (Slide 4):

- Chemical hazards caused by excessive application of a certain food additive can be prevented at the step where the ingredient is added.
- Histamine formation in certain fish species can be prevented at all steps where the proper use of ice, refrigeration, or managing the time out of refrigeration can prevent the product from being exposed to a temperature above 40°F for an extended period of time.
- Chemical hazards such as drug residues in aquaculture products can be prevented at the receiving step by using controls such as supplier declarations or testing.
Principle 2: Determine Critical Control Points

Examples of CCPs where a hazard can be eliminated are (Slide 5):

- Pathogens can be eliminated (killed) at the cooking step by controlling the time and temperature used for cooking.
- Metal fragments that may be in the finished product can be eliminated at a metal detector step because any product containing metal fragments would be removed from the processing line.
- Parasites can be eliminated (killed) at a freezing step by controlling the freezer temperature and how long the product is held at that temperature.

Examples of CCPs where the hazard can be reduced to an acceptable level are (Slide 6):

- The possibility that biological hazards, such as pathogens, and chemical hazards, such as natural toxins, can be reduced to acceptable levels in shellfish at the receiving step if controls are used to ensure that all shellfish are purchased from certified dealers and are properly tagged to document the product has been harvested from approved waters.
- The possibility that unacceptable levels of environmental chemical hazards such as PCBs will be present in fish can be reduced to acceptable levels at the receiving step by ensuring that the fish were not harvested from waters that have been closed by local or state health authorities.
- The possibility of pathogen growth can be reduced to acceptable levels at a storage step by controlling cooler temperatures or using adequate ice.

Multiple CCPs and Multiple Hazards

A single CCP can be used to control more than one hazard. For example, the step where live oysters or clams are received provides an example of when a single CCP could control multiple hazards. Source related hazards for this product such as harvest site pathogens, natural toxins and chemical contaminants could all be controlled at the receiving step by making sure that the shellstock is from an approved source and properly tagged (Slide 7).
In some circumstances, more than one CCP may be needed to control a single significant hazard. For example, receiving and storage steps may be CCPs for a histamine hazard. This is because histamine can develop in these fish at any time if there is time and temperature abuse. Any step in the process where there is a potential for significant time/temperature abuse must be identified as a CCP (Slide 7).

**Multiple Hazards and Single CCP**
- **Product** = Live oysters (shellstock)
- **Hazards** = Harvest site pathogens + Natural Toxins + Chemical Contaminants
- **Single CCP** = Receiving

**Single Hazard and Multiple CCPs**
- **Product** = Fresh Tuna loins
- **Hazard** = Histamine
- **Multiple CCPs** = Receiving + Refrigerated Storage

### CCPs are Product- and Process-Specific
CCPs that have been identified for a product on one processing line may be different for the same product on another processing line. This is because the hazards and their processing controls are impacted by the layout of the plant or processing line, the formulation of the finished product, the process flow diagram or sequence of processing steps, the processing equipment that is used, the sanitation and support programs that are used, and ingredients that may be used (Slide 8).

**Slide 8**

CCP are product- and process-specific and impacted by:
- Layout of the plant or processing line,
- Finished product formulation,
- Process flow or sequence of processing steps,
- Processing equipment,
- Ingredients,
- Sanitation or other support programs.

### Tools to Help Identify CCPs
There are several tools available to help identify which steps are likely to be CCPs for various types of hazards.

**Hazards Guide** – The Hazards Guide provides guidance on likely CCPs for each of the potential seafood safety hazards associated with seafood products. The Hazards Guide has a chapter for each of these food safety hazards. There
is a specific section in each of these chapters that provides information for decisions regarding CCP placement.

**CCP Decision Tree** – Another tool that can help identify which steps are CCPs is the CCP Decision Tree. This tool has a series of questions that can help identify the CCPs in the process. These questions can be applied at each of the processing steps where a significant hazard was identified in the hazard analysis.

The CCP Decision Tree asks a series of three questions that will lead you to decide if a specific processing step is a CCP (Slide 9).

### Slide 9

**CCP Decision Tree**

- **Q 1)** Does this step involve a hazard of sufficient risk and severity to warrant its control?
  - Yes
  - No → Not a CCP

- **Q 2)** Does a control measure for the hazard exist at this step?
  - Yes
  - No → Modify this step, process or product
    - Is control at this step necessary for safety?
      - Yes
      - No → Not a CCP → Stop*

- **Q 3)** Is control at this step necessary to prevent, eliminate or reduce the risk of the hazard to consumers?
  - Yes
  - No → Not a CCP → Stop*

  *Proceed to the next step in process*
Receiving

In the hazard analysis, XYZ Seafood Company identified histamine as a significant food safety hazard at the receiving step. The CCP Decision Tree can be used to determine if the receiving step is a CCP to control the hazard of histamine.

**Question 1)** Does this step involve a hazard of sufficient risk and severity to warrant its control?

**Answer:** Yes – Histamine is a significant hazard for mahi-mahi that could cause consumer illness. Time-temperature abuse during transit could cause histamine to form in fillets.

**Question 2)** Does a control measure for the hazard exist at this step?

**Answer:** Yes – Mahi-mahi fillets are received in containers buried in ice.

**Question 3)** Is control at this step necessary to prevent, eliminate or reduce the risk of the hazard to consumers?

**Answer:** Yes – The hazard of histamine must be prevented at this step.

**Conclusion:** The receiving step is a CCP for the hazard of histamine.

**Example: Fresh Mahi-mahi/XYZ Seafood Co.**

**Complete Hazard Analysis Worksheet**

The hazard analysis worksheet summarizes the results of the complete hazard analysis for XYZ Seafood Company (Slide 10).

XYZ Seafood Company has identified the following CCPs for fresh mahi-mahi fillets:

- Three CCPs to control histamine
  1) Receiving fresh fillets
  2) Refrigerated storage
  3) Finished product refrigerated storage
- One CCP to control food allergens
  1) Weigh/Pack/Label
XYZ Seafood Company – Fresh Mahi-mahi Fillets

<table>
<thead>
<tr>
<th>Processing Step</th>
<th>(2) List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process.</th>
<th>(3) Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No)</th>
<th>(4) Justify the decision that you made in column 3</th>
<th>(5) What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard?</th>
<th>(6) Is this step a Critical Control Point? (Yes or No)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receiving Fresh Fillets</td>
<td>Histamine</td>
<td>Yes</td>
<td>Time/temperature abuse during transit could cause histamine to form in the fish</td>
<td>Mahi-mahi fillets are shipped in containers buried in ice (proper icing)</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step (proper labelling)</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Refrigerated Storage</td>
<td>Histamine</td>
<td>Yes</td>
<td>Time/temperature abuse during storage could cause histamine to form in the fish</td>
<td>Mahi-mahi fillets are shipped in ice &amp; stored in a refrigerated cooler (proper icing)</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step (proper labelling)</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processing Step</td>
<td>(2) List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process.</td>
<td>(3) Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No)</td>
<td>(4) Justify the decision that you made in column 3</td>
<td>(5) What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard?</td>
<td>(6) Is this step a Critical Control Point? (Yes or No)</td>
</tr>
<tr>
<td>-----------------</td>
<td>---------------------------------------------------------------</td>
<td>-------------------------------------------------</td>
<td>-------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>Trim</td>
<td>Histamine</td>
<td>No</td>
<td>Not likely to occur, time at this and weigh/pack/label step is 30 minutes or less</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step (proper labelling)</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Fillet knives are not likely to chip and contaminate product with metal</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weigh/Pack/Label</td>
<td>Histamine</td>
<td>No</td>
<td>Not likely to occur, time at this and weigh/pack/label step is 30 minutes or less</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets are labeled with market name at this step (proper labelling)</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Finished Product Refrigerated Storage</td>
<td>Histamine</td>
<td>Yes</td>
<td>Time/temperature abused could occur during storage</td>
<td>Mahi-mahi fillets are buried in ice &amp; stored in a refrigerated cooler (proper icing)</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>No</td>
<td>Fillets were labeled with market name at Weight/Pack/Label step</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Metal Inclusion</td>
<td>No</td>
<td>Not likely to occur at this step</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
All of the remaining HACCP principles apply only to critical control points (CCPs).

Critical limits must be established for each hazard at each CCP identified in the hazard analysis (Slide 1). This is the third principle of HACCP.

**Slide 1**

In this chapter, you will learn:
- Definition of critical limit.
- How to determine critical limits for a CCP.
- The relationship between critical limits and operating limits.
- Use of the HACCP plan form.

A critical limit represents the boundaries that are used to ensure that a hazard has been controlled (prevented, eliminated, or reduced to an acceptable level) at each CCP (Slide 2). Critical limits must be based on what science or industry experience has demonstrated is necessary to control the hazard.

**Slide 2**

Definition:
Critical Limit: A maximum and/or minimum value to which a biological, chemical or physical parameter must be controlled at a CCP to prevent, eliminate or reduce the occurrence of a food safety hazard to an acceptable level.
Information Sources for Establishing Critical Limits

The Hazards Guide provides critical limit information for most seafood safety hazards. Other information may be needed to establish valid critical limits for a CCP. These other sources of information (see chapter 13) could include process authorities, scientific studies, trade associations, other state or federal regulations. However, in some cases, the appropriate critical limit may not be readily apparent or available. Information may need to be gathered from other sources such as scientific publications, experts or experimental studies (Slide 3).

Sources of Information on Critical Limits (see chapter 13)

<table>
<thead>
<tr>
<th>Information Source</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA Regulations and guidelines</td>
<td>State and local regulations, tolerances and action levels; USDA guidelines, tolerances and action levels, and the National Shellfish Sanitation Program (NSSP) Model Ordinance for Molluscan Shellfish</td>
</tr>
<tr>
<td>Experts</td>
<td>Process authorities; university food scientists/microbiologists, consultants, equipment manufacturers, sanitarians, and trade associations</td>
</tr>
<tr>
<td>Scientific studies</td>
<td>In-house experiments and contract labs or universities</td>
</tr>
<tr>
<td>Scientific information</td>
<td>Journal articles, food science texts, microbiology texts, and National Seafood HACCP Alliance Compendium</td>
</tr>
</tbody>
</table>

There are many different types of critical limits. They must be specific for the critical control point and the hazard that is being controlled (Slide 4 and 5). Different critical limits may be needed for species-related hazards and process-related hazards.

Each CCP must have one or more critical limits for each food-safety hazard (Slide 4). An effective critical limit will define what can be measured or observed to demonstrate that the hazard is being controlled at that CCP. For example, both time and temperature are necessary elements of a critical limit to eliminate food safety hazards like pathogens at a cook step.
### Slide 4

**Examples of Critical Limits for species-related hazards**

<table>
<thead>
<tr>
<th>Product</th>
<th>Significant Hazard</th>
<th>Critical Control Point</th>
<th>Critical Limits</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aquacultured shrimp</td>
<td>Aquaculture drugs</td>
<td>Receiving (from farm)</td>
<td>Suppliers certificate on file (indicating proper drug use)</td>
</tr>
<tr>
<td>Oysters (live)</td>
<td>Natural toxins</td>
<td>Receiving (from harvester)</td>
<td>All shellstock tagged with the date and place of harvest, type and quantity of shellfish, and name or registration number of harvest vessel and all shellstock from waters approved by State Shellfish Authority and all shellstock from a licensed harvester</td>
</tr>
<tr>
<td>Raw Tuna</td>
<td>Histamine</td>
<td>Storage</td>
<td>Fish are completely surrounded by ice</td>
</tr>
</tbody>
</table>

### Slide 5

**Examples of Critical Limits for process-related hazards**

<table>
<thead>
<tr>
<th>Product</th>
<th>Significant Hazard</th>
<th>Critical Control Point</th>
<th>Critical Limits</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battered fish</td>
<td><em>Staphylococcus aureus</em> growth and toxin formation</td>
<td>Batter application</td>
<td>Hydrated batter does not exceed 50°F for more than 12 hrs. or 70°F for more than 3 hrs., cumulative.</td>
</tr>
<tr>
<td>Imitation crabmeat</td>
<td>Metal inclusion</td>
<td>Metal detector (after packaging)</td>
<td>No detectable metal fragments in finished product</td>
</tr>
<tr>
<td>Hot smoked fish, vacuum packaged</td>
<td><em>Clostridium botulinum</em> toxin formation (in finished product)</td>
<td>Hot smoking</td>
<td>Internal fish temperature held at or above 145°F for at least 30 minutes</td>
</tr>
<tr>
<td>Ready-to-eat seafood salad</td>
<td>Pathogen growth</td>
<td>Cooler storage</td>
<td>Cooler temperature not to exceed 40°F</td>
</tr>
</tbody>
</table>
Critical Limit Options

Processors may have different options for controlling a particular hazard. Each control option usually requires the use of different critical limits. The selection of the best control option and the best critical limit is often driven by practicality and experience. The following examples describe three different options for effective control measures and critical limits that could be applied at a fryer (cooking) CCP to eliminate the hazard of bacterial pathogens in fried fish cakes.

Option 1 is not typically the best option (Slide 7). Setting a critical limit such as “no pathogens detected” is rarely appropriate. This type of critical limit is difficult to monitor, and testing to determine critical limit deviations may require several days. Critical limits must allow monitoring on a timely basis. Sampling and testing is normally more appropriate as a verification step, described later in this course.

<table>
<thead>
<tr>
<th>Hazard</th>
<th>CCP</th>
<th>Critical Limits</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pathogen survival through cooking</td>
<td>Cooker</td>
<td>&gt;160°F internal product temperature for ≥1.5 minutes for elimination of pathogens of concern in cooked crabs (e.g. <em>Listeria monocytogenes</em>)</td>
</tr>
<tr>
<td>Pathogen growth</td>
<td>Drying oven</td>
<td>Drying schedule — oven temperature: ≥ 200°F, time ≥120 min., air flow rate: ≥ 2 ft³/min, product thickness ≤0.5 inches (to achieve aw of 0.85 to control pathogens in dried foods)</td>
</tr>
<tr>
<td>Pathogen growth</td>
<td>Acidification</td>
<td>Batch schedule — product weight, ≤ 100 lbs.; soak time, ≥ 8 hrs; acetic acid concentration, ≥ 3.5 percent; volume ≤ 50 gal. (to achieve maximum pH of 4.6 to control <em>Clostridium botulinum</em> in pickled foods)</td>
</tr>
</tbody>
</table>
Option 2 uses the internal product temperature and time achieved during frying as a critical limit (Slide 8). This critical limit option is more practical than finished product pathogen testing. However, internal product temperature and time cannot be easily monitored for all of the products that are cooked, and heat transfer rates during cooking could vary for a variety of reasons. For this reason, it would be difficult to measure whether or not this critical limit has been met for all products.

Slide 8
Option No. 2
Product: Fish cakes
Hazard — pathogen survival through cooking
CCP — fryer
Critical limit — minimum internal temperature of 165°F for 36 seconds

It seldom is practical to continually monitor the internal temperature of the food product to ensure conformance with a critical limit. As an alternative, critical limits such as those in Option 3 may establish conditions necessary to ensure that the cooking process achieves the minimum product temperature and time. In this option, the oil temperature, the fish cake thickness and the time that the cake stays in the hot oil are all factors that affect the final fish cake temperature (Slide 9). These factors are easy to monitor and measurements are obtained quickly to determine that critical limits have been met. A scientific study (validation) must be performed to ensure that controlling these factors will always result in an internal product temperature that will destroy pathogens of concern. Typically, this option is better than the two previous options even though more critical limits are involved.

Slide 9
Option No. 3
Product: Fish cakes
Hazard — pathogen survival
CCP — fryer
Critical limit — minimum fryer oil temperature of 350°F
Critical limit — maximum fish cake thickness of ¾ inch
Critical limit — minimum cook time in the oil of two minutes

Operating Limits and Critical Limits

An operating limit allows the detection of a potential problem before a critical limit is violated (Slide 10). Operating limits should not be confused with critical limits. Operating limits are established at a level that would be reached before the critical limit is violated. The process should be adjusted when the operating limit is reached to avoid violating critical limits. These actions are called
process adjustments. A processor may use these adjustments to avoid loss of control and the need to take corrective action. Spotting a trend toward loss of control early and acting on it can save product re-work, or worse yet, product destruction.

**Slide 10**

**Definition:**
Operating Limits: Criteria that are more stringent than critical limits and that are used by an operator to reduce the risk of a deviation.

Operating limits may be selected for various reasons:

- For quality reasons (e.g., higher cooking temperatures may enhance flavor development or to control organisms that can cause spoilage);
- To avoid deviating from a Critical Limit, processors often establish an Operating Limit that is more stringent that the Critical Limit. For example, a processor could establish a cooking temperature Operating Limit that is higher than the HACCP Critical Limit. If monitoring indicated that temperatures fell below the Operating Limit, the processor would have time to initiate a Process Adjustment to avoid a Critical Limit deviation;
- To account for normal variability (e.g., a fryer with a $5^\circ$F variability should be set at least $5^\circ$F above the critical limit to avoid violating it).

Slide 11 illustrates several important points:

- Operating limits and process adjustments,
- Critical limits and corrective actions, and
- Importance of lot size.

In this example, of a cooking process, an operating limit is established at $200^\circ$F and a critical limit at $190^\circ$F. Somewhere in the $10^\circ$F range between these two points, processors will make a process adjustment to bring the cook temperature back above $200^\circ$F. Because an adjustment is made before the temperature drops below the critical limit of $190^\circ$F, no corrective action record is required. However, if an adjustment is not taken until after the temperature drops below the critical limit, as shown in Slide 11, appropriate corrective actions must be taken and a corrective action report must be placed in the HACCP records file (corrective actions and records will be discussed in subsequent chapters).

When a corrective action is necessary, processors must be able to identify and segregate the affected lots. If lot sizes are big (Figure 1), large quantities of product may require segregation and corrective action despite the fact that only a small amount of product was produced when critical limits were exceeded. Lot size also relates to effective traceability and recalls. Coding production into smaller lots (Figure 2) means less product may be involved when violation of a critical limit occurs. Therefore, processors should change codes often and match monitoring frequency with code changes.
HACCP Plan Form

A standardized HACCP plan form (Slide 12) is designed to ensure that HACCP Principles 3 through 7 are adequately described when the HACCP plan is developed. A written HACCP plan is required by the FDA Seafood HACCP Regulation when the hazard analysis determines that there are one or more significant safety hazards, and this plan must be made available to inspectors.
The HACCP plan form is used to:

- List the CCPs from the hazard analysis worksheet in the first column.
- List the significant hazards at each CCP in the second column.
- List the critical limits for each significant hazard in the third column.
- List all elements of monitoring (what, how, frequency, and who) in the fourth column.
- List corrective actions in the fifth column.
- List verification procedures in sixth column.
- List records in the last column.

**Example: Fresh Mahi-mahi/XYZ Seafood Co.**

XYZ Seafood Company’s hazard analysis, described in the previous two chapters, identified four critical controls points, including: 1) the receiving step is a CCP for the hazard of histamine; 2) the refrigerated storage step is a CCP for the hazard of histamine; 3) the weigh/pack/label step is a CCP for the hazard of food allergens; and 4) the finished product storage step is a CCP for the hazard of histamine.

**Receiving CCP:** The HACCP team used the Hazards Guide to determine the critical limits for this CCP. XYZ Seafood Company is a secondary processor who receives and stores mahi-mahi fillets on ice. Control Strategy 3, Transit Control, in Chapter 7 (Histamine) of the Hazards Guide is the best control strategy. This strategy recommends a critical limit of: *fish completely surrounded by ice at the time of delivery.*

This critical limit is entered in the HACCP plan form.

**Refrigerated Storage** is the second CCP for the hazard of histamine. This step and the **Finished Product Storage** CCP occur in the same cooler. Both steps also have same hazard (histamine), and the same control strategy is used for each step. The HACCP Team identified Storage Controls in Chapter 7 (Histamine) of the Hazards Guide as the best control strategy for their CCP. This strategy recommends a critical limit of: *Mahi-mahi fillets are completely surrounded by ice throughout storage time.*

This critical limit is entered in the HACCP plan form for each of these two CCPs.

**Weigh/Pack/Label CCP:** The HACCP Team identified Control Strategy eight. Finished Product Labeling Controls, in Chapter 19 (Major Food Allergens and Food Additives) of the Hazards Guide as the best control strategy for their situation. This strategy recommends a critical limit of: *all finished product containers will be labeled with the correct market name of the fish.*

This critical limit is entered in the HACCP plan form for this CCP.
Complete HACCP Plan Form – Critical Limits

The HACCP plan form must list the critical limits at each CCP. The first three columns of the HACCP plan form have been completed for XYZ Seafood Company’s fresh mahi-mahi fillets (Slide 13).
### HACCP Plan Form

<table>
<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring</th>
<th>Corrective Action</th>
<th>Verification</th>
<th>Records</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Mahi-mahi fillets are completely surrounded with ice at receipt.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Food Allergens</td>
<td>All finished product containers will be labeled with the correct market name of the fish.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Firm Name:** XYZ Seafood Company

**Product:** Fresh mahi-mahi fillets

**Firm Address:**
238 Coastal Lane, Happy Beach, XX

**Method of Storage and Distribution:** Stored and distributed buried in ice

**Intended Use and Consumer:** To be cooked and consumed by the general public

**Signature:**

**Print name:**

**Date:**
Chapter 8

Principle 4: Critical Control Point Monitoring

CCP monitoring is used to ensure that a critical limit is met (Slide 1). Monitoring is the fourth principle of HACCP.

Slide 1

In this chapter, you will learn:
• Definition of monitoring,
• Purpose of monitoring,
• Design of a monitoring system,
• Methods and equipment for monitoring critical limits.

Monitoring involves the selection of appropriate measurements or observations at a specified frequency to ensure that a CCP is under control (Slide 2).

Slide 2

Definition:
Monitoring: A planned sequence of observations or measurements to assess whether a CCP is under control and to produce an accurate record to demonstrate that critical limits have been met.

The purpose of monitoring is to ensure that the critical limit has been met and the food safety hazard is being controlled. Monitoring also provides data for records to document that products were produced in compliance with the HACCP plan. It is important that monitoring procedures are specific for the identified critical limit. For example, if the critical limit requires adequate ice,
the monitoring procedure would be a visual check for ice and not a check for product temperature. When a critical limit is not met, a corrective action is needed (Slide 3).

**Slide 3**

Purpose of Monitoring:
- To ensure that a critical limit is met,
- To provide documentation that critical limits have been met,
- To identify when there is loss of control (a deviation occurs at a CCP).

There are four elements that are required in an effective monitoring system (Slide 4).

**Slide 4**

Elements of Monitoring
- What will be monitored?
- How will monitoring be performed?
- What is the frequency of monitoring?
- Who will conduct the monitoring?

**What will be monitored?**

Monitoring may involve measuring a characteristic of the product or process to determine if a critical limit is met at a CCP (Slide 5).

Examples of monitoring **measurements** could include:

- Cold-storage temperature when cooler temperature is part of the critical limit.
- The pH of an acidifying ingredient when pH is part of the critical limit.
- Line speed and cooker temperature when cook time and temperature are part of the critical limit.

Monitoring may also involve observations to determine if a critical limit is met at a CCP.

Examples of monitoring **observations** could include:

- Look for a vendor’s certificate that accompanies a lot of raw material when approved source is part of the critical limit.
- Check if fish are surrounded with ice when adequacy of ice is part of the critical limit.
- Check to see that harvest area is listed on a tag attached to a container of molluscan shellfish when approved source is part of the critical limit.
Principle 4: Critical Control Point Monitoring

Slide 5

What will be monitored?
A measurement or observation to assess if the CCP is operating within the critical limit.

How will monitoring be performed?

Different methods can be used to monitor critical limits (Slide 6). These methods need to be real-time and accurate.

Using instruments to measure a critical limit quantity is an effective way to conduct monitoring at a CCP. Examples of monitoring instruments could include thermometers, pH meters, water activity meters, data loggers, etc.

Monitoring methods can also involve visually checking what you are monitoring. For example, a visual check for the adequacy of ice; an evaluation of sensory attributes of products; or a visual check of supplier certificates.

It must be clear from an observation whether or not a Critical Limit has been violated. For example, a Critical Limit of “adequate use of ice” is subjective and imprecise, making it difficult to monitor. Monitoring and records must be unambiguous and can be acted upon. Monitoring should be designed to provide rapid, real-time results. Microbiological testing is seldom effective for monitoring CCPs. Very often the analytical methods are lengthy and large sample sizes are usually needed to ensure all units of product conform to microbiological limits. There is no time for lengthy analytical testing during routine monitoring because critical limit failures must be detected quickly and an appropriate corrective action instituted before product shipment.

Slide 6

How will monitoring be performed?

• Measurements (quantitative critical limits) or observations (qualitative critical limits).
• Needs to be real-time and accurate.

What is the frequency of monitoring?

Monitoring frequency will depend on the critical limit and the types of observations and measurements that are needed. The frequency of monitoring can be at regularly scheduled intervals (non-continuous) or continuous (Slide 7).

Slide 7

What is the frequency of monitoring?

• Monitoring frequency should be sufficient to ensure that the critical limit is met.
• Monitoring frequency can be non-continuous or continuous.
Non-continuous Monitoring

It is necessary to establish a monitoring interval that ensures critical limits are met. The frequency of non-continuous (periodic) monitoring could be influenced by historical knowledge of the product and process.

Questions that could help determine the correct frequency include:

- How much does the process normally vary (e.g., how consistent is the data)? If the monitoring data shows a great deal of variation, the time between monitoring checks should be short.
- How close are the normal operating values to the critical limit? If the normal values are close to the critical limit, the time between monitoring checks should be short.
- How much product is the processor prepared to risk if the critical limit is exceeded?

Examples of non-continuous monitoring include:

- Visual daily checks that fish are adequately iced.
- Sensory examination for decomposition in histamine-forming seafood at receipt from the vessel.
- End Point Internal Product Temperature (EPIPT) of cooked crab sections.

Continuous Monitoring

When possible, continuous monitoring procedures should be used. Continuous monitoring is generally performed by an instrument that produces a continuous record. The record needs to be checked periodically to ensure that the critical limit is being met. The length of time between checks will directly affect the amount of rework or product loss when a critical limit deviation is found.

Examples of continuous monitoring could include:

- The time and temperature of a batch pasteurization process for crabmeat may be continuously monitored and recorded on a temperature-recording chart.
- The temperature of a molluscan shellfish storage cooler is continuously monitored and recorded.

Who will Monitor?

Assignment of the responsibility for monitoring is an important consideration when developing a HACCP plan (Slide 8).
Individuals assigned to CCP monitoring can be:

- Line personnel,
- Equipment operators,
- Supervisors,
- Maintenance personnel, or
- Quality assurance personnel.

Monitoring by line personnel and equipment operators can be advantageous since they are continuously watching the product and/or equipment. Including production workers in HACCP activities has the advantage of building a broad base of understanding and commitment to the HACCP program.

The monitor’s duties should require that all deviations from operating limits and critical limits be reported immediately to make sure process adjustments and corrective actions are made in a timely manner. All records and documents associated with CCP monitoring must be signed or initialed by the person doing the monitoring (Slide 9).

**Slide 9**

Those responsible for monitoring a CCP should:
- Be trained in the CCP monitoring techniques.
- Fully understand the importance of CCP monitoring.
- Have ready access to the monitoring activity.
- Accurately report each monitoring activity.
- Immediately report critical limit deviations.

Properly trained personnel must be available at all times that the CCP requires monitoring. Additional monitoring personnel may be needed when monitoring is required during breaks, weekends, or when monitoring is required throughout multiple work shifts.

Some examples of measurements and observations that could be used for monitoring a critical limit at a CCP are provided below (Slide 10).

**Slide 10**

Monitoring Examples:
- Time and temperature of process
- Time and internal temperature combinations
- Water activity (a_w)
- pH
- Internal product temperature
- Salt concentration in brine
- Metal inclusion screening
Critical Limit for a Cooking CCP: minimum cooker temperature of 212°F for a minimum of three minutes

- What will be monitored? cooker temperature and time
- How will it be monitored? time/temperature recording device and visual observation
- Frequency of monitoring: continuous with visual check for each batch
- Who will monitor? cooker/operator

Critical Limit for Refrigerated Storage Step: all fish completely surrounded by ice

- What will be monitored? adequacy of ice
- How will it be monitored? visual
- Frequency of monitoring: twice a day
- Who will monitor? cooler person

Critical Limit at Acidification Step: acidity (pH) less than 4.6

- What will be monitored? pH
- How will it be monitored? pH meter
- Frequency of monitoring: each batch
- Who will monitor? QC person

Critical Limit at Labeling Step: product label identifies market name of seafood

- What will be monitored? finished product label
- How will it be monitored? visual
- Frequency of monitoring: representative number of samples from each lot
- Who will monitor? packaging manager

Monitoring Equipment

The selection of the proper monitoring equipment, which could include instruments such as thermometers and rapid test kits, is a major consideration during development of a HACCP plan. There are many different kinds of monitoring instruments or tools that may be appropriate for different critical limits (Slide 11).

Examples of monitoring equipment could include:
- thermometers
- recorder charts
- clocks
- pH meters
- water activity meters
- data loggers
- metal detectors
- salometer
Principle 4: Critical Control Point Monitoring

The equipment chosen for monitoring at the CCP must be accurate to ensure control of the hazard. Equipment used to monitor critical limits must be calibrated at a frequency that ensures accuracy. For example, the accuracy of temperature measuring instruments should be checked frequently or as recommended by the device manufacturer. Accuracy is further discussed in Chapter 10 (Principle 6: Establish Verification Procedures).

Example: Fresh Mahi-mahi/XYZ Seafood Co.

XYZ Seafood Company identified four critical controls points in the hazard analysis. There are three CCPs for the hazard of histamine: receiving, refrigerated storage, and finished product storage. There is one CCP for the hazard of food allergens: weigh/pack/label.

**Receiving CCP:** The HACCP team used the Hazards Guide to determine the monitoring procedures for the CCP. The HACCP Team identified Control Strategy 3, Transit Control, in Chapter 7 (Histamine) of the Hazards Guide as the best control strategy for their situation. This strategy recommends a monitoring procedure that includes the following elements:

- **What:** adequacy of ice surrounding containers of fillets at delivery
- **How:** visual check of adequacy of ice in a representative number of containers at delivery
- **Frequency:** every delivery
- **Who:** receiving manager

This monitoring information is entered in the HACCP plan form.

**Refrigerated Storage and Finished Product Storage CCPs:** The HACCP team identified “Control Strategy 5, Storage Control,” in Chapter 7 (Histamine) of the Hazards Guide as the best control strategy for their CCP. This strategy recommends a monitoring procedure that includes the following elements:

- **What:** adequacy of ice surrounding containers of fillets in the cooler
- **How:** visual check of adequacy of ice in a representative number of containers in the cooler
- **Frequency:** at the beginning and end of the work day
- **Who:** cooler manager

This monitoring information is entered in the HACCP plan form.

**Note**

Representative numbers can be based on the size of the lot or total number of containers and experience regarding the possible amount of variation within the lot.
**Weigh/Pack/Label CCP:** The HACCP team identified “Control Strategy 8, Finished Product Labeling Controls,” in Chapter 19 (Major Food Allergens and Food Additives) of the Hazards Guide as the best control strategy for their situation. This strategy recommends a monitoring procedure that includes the following elements:

- **What:** The market name on each container of finished product
- **How:** Visual check finished product label
- **Frequency:** Representative number of finished product containers per lot
- **Who:** Packing manager

This monitoring information is entered in the HACCP plan form.

**Example: Fresh Mahi-mahi/XYZ Seafood Co.**

**Complete HACCP Plan Form – Monitoring**

The HACCP plan form must list the monitoring procedures at each CCP (Slide 12).
## HACCP Plan Form

**Firm Name:** XYZ Seafood Company

**Product:** Fresh mahi-mahi fillets

### Critical Control Point (CCP)

<table>
<thead>
<tr>
<th>CCP</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring What</th>
<th>How</th>
<th>Frequency</th>
<th>Who</th>
</tr>
</thead>
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<td>Mahi-mahi fillets are completely surrounded with ice at receipt.</td>
<td>Adequacy of ice surrounding mahi-mahi fillets at delivery</td>
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<td>Every Delivery</td>
<td>Receiving Manager</td>
</tr>
<tr>
<td>Refrigerated Storage</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td>Adequacy of ice surrounding mahi-mahi fillets</td>
<td>Visual check of adequacy of ice in a representative number of containers in cooler storage</td>
<td>At the beginning and end of the work day</td>
<td>Cooler Manager</td>
</tr>
<tr>
<td>Weigh/Pack/Label</td>
<td>Food Allergens</td>
<td>All finished product containers will be labeled with the correct market name.</td>
<td>The market name on each container of finished product</td>
<td>Visual check of a representative number of containers and their label</td>
<td>Each customer order</td>
<td>Packing Manager</td>
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<td>Finished Product Refrigerated Storage</td>
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<td>Visual check of representative number of containers in cooler storage</td>
<td>At the beginning and end of the work day</td>
<td>Cooler Manager</td>
</tr>
</tbody>
</table>

### Firm Name:

XYZ Seafood Company

**Firm Address:**
238 Coastal Lane, Happy Beach, XX

**Method of Storage and Distribution:** Stored and distributed buried in ice

**Intended Use and Consumer:** To be cooked and consumed by the general public

**Signature:**

**Date:**

**Print name:**
Chapter 9

Principle 5: Corrective Actions

Corrective Actions are taken when a critical limit is not met. Corrective action is the fifth principle of HACCP (Slide 1).

Slide 1

In this chapter, you will learn:
• The definition of corrective actions,
• Procedures for corrective actions, and
• Record-keeping requirements for corrective actions.

A HACCP system should be designed to ensure that critical limit deviations are rapidly identified and corrected (Slide 2). The responsibility for taking corrective actions must be assigned to one or more individuals who have a thorough understanding of the operation, the products, the firm’s HACCP plan, and the authority to make decisions.

Slide 2

Definition:
Corrective Action: Procedures to be followed when a deviation occurs.

Predetermined Corrective Actions

Predetermined corrective actions provide a “how-to” guide that describes the steps that need to be taken when a critical limit deviation occurs (Slide 3). It
may be possible, and is always desirable, to correct the problem immediately. Although it may not be possible to anticipate all the deviations that could happen, corrective actions still need to be taken even when an unanticipated situation occurs.

Components of Corrective Actions

There are two essential parts to a corrective action (Slide 4). The objectives of the corrective action are to keep potentially unsafe product from reaching the consumer and to restore control to the process prior to producing more product.

When a deviation is detected, the first action is to identify the product involved. This product should be segregated and evaluated to determine if a food safety hazard exists. If a hazard exists, the affected product must be reworked or destroyed to ensure it will not cause consumer illness.

Process control must also be restored. Corrective actions must bring the CCP back under control. A corrective action should take care of the immediate (short-term) problem as well as provide long-term solutions. The objective is to re-establish control of the process so that production can start again as soon as possible without further deviations.

Tools to Help Evaluate Product Safety

A qualified individual must be assigned responsibility for evaluating product safety (Slide 5). Not every firm has an expert on staff that can evaluate the safety of products involved in a deviation. It may be necessary to identify additional resources that can help with product safety evaluations.

The Hazards Guide is an important tool that can assist with corrective actions and any necessary product safety evaluations. For example, Appendix 4 of the Hazards Guide provides information on bacterial pathogen growth and inactivation. Hazards Guide Table #A-2 provides maximum cumulative time/temperature guidance for controlling pathogen growth and toxin formation in seafood. Information on appropriate corrective actions is also outlined in
the control strategies provided in each of the hazard specific chapters in the Hazards Guide.

### Slide 5

**Tools to help evaluate product safety:**
- Food Safety Experts
- Production monitoring data/records
- NSSP Shellfish Model Ordinance
- Hazards Guide
  - Appendix 4: Pathogen Tables
  - Appendix 5: Guidance Levels
- Laboratory testing

### Examples:

- Using the Hazards Guide, a processor who produces a battered seafood product has determined that the batter step is a CCP for the hazard of *Staphylococcus aureus* toxin. The critical limit is: batter temperature is at or below 50°F. If monitoring shows that the batter temperature is 65°F, the critical limit has not been met and a corrective action must be taken. To help evaluate product safety, Hazards Guide Table #A-2 can be used to determine if toxin production could have occurred. This table shows that cumulative exposure time to temperatures between 50°F and 70°F must be no more than 12 hours for toxin to be produced. The monitoring records show that the exposure time above 50°F and less than 70°F was less than 4 hours. Therefore, the product produced with this batter was not exposed to conditions that would have allowed toxin to be produced and created a food safety hazard.

- For bivalve molluscan shellfish operations, the NSSP Shellfish Model Ordinance ([www.issc.org](http://www.issc.org)) provides information that may be used to evaluate product safety when a deviation occurs.

- Laboratory testing could also provide valuable information to evaluate product safety. Important considerations when using test results include: a valid sampling scheme must be used, the pertinent pathogen or chemical of interest must be accurately identified, and an approved/recognized testing method must be used. If a product is to be tested and released, the sampling method is very important. The use of a faulty sampling protocol can result in accepting, rather than rejecting, an undesirable product. The limits of sampling plans must be understood. It may be prudent to consult an expert.

### Determine Product Disposition

A proper and thorough safety evaluation is necessary to determine the disposition of the product (Slide 6). It is best to be cautious, but product destruction may not always be necessary. Decisions related to the disposition of the affected product must be based on sound evidence that the deviation did not create a food safety hazard. This evidence must be documented to
support the decision. Like other Corrective Actions, if the product is rejected or destroyed, the processor needs to document that this has been done.

### Slide 6

**Steps to determine the disposition of product:**

**Step 1:** Determine if the product presents a safety hazard.

**Step 2:** If no hazard exists, the product may be released.

**Step 3:** If a potential hazard exists, determine if the product can be:
   a) Reworked/reprocessed, or
   b) Diverted for a safe use.

**Step 4:** If a food safety hazard does exist, the product must be rejected or destroyed.

### Correct and Eliminate the Cause of the Deviation and Restore Process Control

It is necessary to determine the cause of the deviation to prevent the same problems from occurring again (Slide 7). **When critical limit deviations frequently reoccur, the process and the HACCP plan must be re-evaluated.** The initial cause or causes of the process deviation must be identified and corrected so that process control can be restored.

### Slide 7

Corrective actions must identify the cause of the deviation and restore process control.

### Documenting Corrective Actions

Corrective actions must be documented (Slide 8). Corrective action records will show how the safety of the product was evaluated and its disposition. This record will also document the actions taken to fix the problem that caused the deviation and restore process control. These records will help the firm identify recurring problems. This information can be used to evaluate and modify the HACCP plan, if necessary.

### Slide 8

Corrective actions must be documented to indicate the safety status and consequences for the products and process involved.

Processors must develop a corrective-action record to ensure that all of the necessary information is documented (Slide 9). This corrective action report is required by the FDA Seafood HACCP regulation.
A corrective action record would typically include:

a) Product identification (product description and the amount on hold)
b) Description of the deviation, including time and date of deviation
c) Results of the food safety evaluation including test results, when necessary
d) Corrective action taken including:
   • How the cause of the deviation was corrected
   • Final disposition of the affected product
e) Name and signature of the individual responsible for taking the corrective action
f) Signature and date of the review
Corrective Action Examples

Slides 10 and 11 provide examples for simple corrective actions for various types of food safety hazards. They are best written in an “If/Then” format. The “If” part of the corrective action describes the deviation condition, and the “then” part describes the action taken.

### Slide 10

**Corrective action examples for species-related hazards**

<table>
<thead>
<tr>
<th>Critical Control Point</th>
<th>Significant Hazard</th>
<th>Critical Limit</th>
<th>Corrective Actions</th>
</tr>
</thead>
</table>
| Receiving aquacultured shrimp from the farm | Aquaculture drugs | Supplier certificate on file (indicating proper drug use) | If: supplier certificate is not on file;  
Then: reject lot and discontinue using supplier until appropriate, accurate certificate obtained. |
| Receiving live oysters from the harvester | Natural toxins | All shellstock tagged with the date and place of harvest, type and quantity of shellfish, and name or registration number of harvest vessel and All shellstock from waters approved by State Shellfish Authority and All shellstock from a licensed harvester. | If: shellstock tags are missing and/or do not have required information;  
Then: reject shellstock.  
If: harvester not licensed or harvest waters are not approved;  
Then: reject shellstock and discontinue purchasing from harvester until properly licensed. |
Corrective action examples for process-related hazards

<table>
<thead>
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<th>Significant Hazard</th>
<th>Critical Limit</th>
<th>Corrective Actions</th>
</tr>
</thead>
</table>
| Batter application                      | *Staphylococcus aureus* growth and toxin formation | Hydrated batter does not exceed 50°F for more than 12 hrs. or 70°F for more than 3 hrs., cumulatively | **If:** batter temperature and time (cumulative) exceeds critical limits;  
**Then:** destroy batter and product produced during period of deviation or hold and evaluate product for product safety, and adjust/repair refrigeration equipment for batter. |
| Metal detector (after packaging)       | Metal inclusion                              | No detectable metal fragments in product                                        | **If:** product is rejected by metal detector;  
**Then:** rework product to remove metal if possible and pass through metal detector or destroy product, and re-calibrate metal detector to determine if it is working properly and adjust as necessary and determine the source of metal and fix the problem. |
| Hot smoking (vacuum packaged)          | *Clostridium botulinum* toxin formation (in finished product) | Internal fish temperature held at or above 145°F for at least 30 minutes       | **If:** product does not reach required internal temperature for the required time;  
**Then:** extend cook time until proper internal temperature is met or re-cook product to 145°F for 30 minutes or destroy product, and make repairs/adjustments to equipment to ensure process meets critical limits. |
Critical Control Point | Significant Hazard | Critical Limit | Corrective Actions
--- | --- | --- | ---
Cooler storage (Ready-to-eat seafood salad) | Pathogen growth | Cooler temperature not to exceed 40°F | If: cooler temperature goes above 40°F; Then: chill and hold affected product and determine cumulative time of temperature deviation. Use Appendix 4 in Hazards Guide to determine if safety hazard exists for pertinent pathogen or destroy product, and make repairs/adjustments to cooler.

Example: Fresh Mahi-mahi/XYZ Seafood Co.

XYZ Seafood Company identified four critical controls points in the hazard analysis. There are three CCPs for the hazard of histamine: receiving, refrigerated storage and finished product storage. There is one CCP for the hazard of food allergens: weigh/pack/label. The HACCP team used the Hazards Guide to determine what corrective action procedures are needed in their HACCP plan.

**Receiving CCP:** The HACCP Team identified “Control Strategy 3, Transit Control,” in Chapter 7 (Histamine) of the Hazards Guide as the best control strategy for their situation. This control strategy recommends a corrective action of:

*If:* the amount of ice is not adequate;

*Then:* reject product. Call supplier to let them know critical limit was not met and provide product delivery specifications. Discontinue use of supplier until their transport procedures are corrected.

This corrective action is entered in the HACCP plan form.

**Note**

In some instances where there is historical information on the transport of the product, the Corrective Action could state:

*If:* the amount of ice is not adequate; *Then:* check the internal temperature of the exposed fish. Reject all fish that exceed 40°F internal temperature or lack sufficient transport history.
Refrigerated Storage and Finished Product Storage CCPs: The HACCP Team identified Storage Controls in Chapter 7 (Histamine) of the Hazards Guide as the best control strategy for their CCP. This strategy recommends a corrective action of:

**If:** the amount of ice is not adequate;

**Then:** Add ice and hold the product until it can be evaluated based on its total time and temperature exposure, including exposures during prior processing operations. Determine the cause of the problem and fix it.

This corrective action is entered in the HACCP plan form.

Weigh/Pack/Label CCP: The HACCP Team identified Finished Product Labeling Controls in Chapter 19 (Major Food Allergens and Food Additives) of the Hazards Guide as the best control strategy for their situation. This strategy recommends a corrective action of:

**If:** a container is improperly labeled,

**Then:** segregate it and properly label it before the customer order is placed in the finished product cooler. Modify labeling procedure and conduct training as necessary to ensure that all products are properly identified.

This corrective action is entered in the HACCP plan form.

Example: Fresh Mahi-mahi/XYZ Seafood Co.

**Complete HACCP Plan Form – Corrective Action**

Slide 12 HACCP plan form for XYZ Seafood Company completed through Corrective Action
HACCP plan form for XYZ Seafood Company completed through corrective action

<table>
<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring</th>
<th>Corrective Action</th>
<th>Verification</th>
<th>Records</th>
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<td>Adequacy of ice surrounding mahi-mahi fillets at delivery</td>
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<td>Refrigerated Storage</td>
<td>Histamine</td>
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<td>Visual check of adequacy of ice in a representative number of containers in cooler storage</td>
<td>At the beginning and end of the work day</td>
<td>Cooler Manager</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>If: the amount of ice is not adequate. Then: chill and hold the product until it can be evaluated based on its total time and temperature exposure, including exposures during prior processing operations, and add ice and make adjustments to the ice application process.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weigh/Pack/Label</td>
<td>Food Allergens</td>
<td>All finished product containers will be labeled with the correct market name of the fish.</td>
<td>The market name on each container of finished product</td>
<td>Visual check of a representative number of containers and their label</td>
<td>Each customer order</td>
<td>Packing Manager</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>If: a container is improperly labeled. Then: segregate it and properly label it before the customer order is placed in the finished product cooler, and modify labeling procedure and conduct training as necessary to ensure that all products are properly identified.</td>
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**Firm Name:** XYZ Seafood Company

**Product:** Fresh mahi-mahi fillets

**Firm Address:**
238 Coastal Lane, Happy Beach, XX

**Method of Storage and Distribution:** Stored and distributed buried in ice

**Intended Use and Consumer:** To be cooked and consumed by the general public

**Signature:**

**Print name:**

**Date:**
Chapter 10

Principle 6: Establish Verification Procedures

The sixth principle of HACCP requires that verification procedures be established to assure the HACCP program is effective (Slide 1).

In this module, you will learn:
- The definition of verification
- Validation as part of verification
- Verification procedures

Verification includes the procedures that are needed to ensure that the HACCP plan is designed properly and has been implemented correctly (Slide 2). The validity of the HACCP plan is determined before the plan is implemented. Routine verification is used to determine if the plan is working properly.

Verification

The purpose of the overall HACCP plan is to control food safety hazards. The purpose of verification is to provide a level of confidence that the plan is based
on solid scientific principles, is adequate to control the hazards associated with
the product and process, and is being followed. A key concept in the verification
principle is “Trust what you can verify” (Slide 3).

Verification can be complex because there are several elements associated with
this principle (Slide 4). The types of verification activities that may be needed
include: validation, CCP verification, HACCP system verification, and regulatory
verification. Each processor must determine which activities are necessary for
their unique situation.

Slide 3

“Trust what you can verify.”

Slide 4

Types of Verification Procedures:
1) Validation (before the HACCP plan is implemented)
2) CCP verification (regularly scheduled activities):
   • Calibration of process-monitoring devices,
   • Record review,
   • Targeted sampling and testing.
3) HACCP system verification (periodic activity):
   • HACCP plan reassessment
   • Microbiological end-product testing and third party audits
4) Regulatory verification (periodic activity)

1) Validation

Validation is an essential component of verification and requires substantiation
that the HACCP plan, if implemented effectively, is sufficient to control the
significant food-safety hazards (Slide 5). Validation of the plan occurs before the
plan is actually implemented. The purpose of validation is to provide objective
evidence that all essential elements of the plan have a scientific basis and
represent a “valid” approach to controlling the food-safety hazards associated
with a specific product and process.

Slide 5

Definition:
Validation: The element of verification focused on collecting and
evaluating scientific and technical information to determine if the
HACCP plan, when properly implemented, will effectively control the
hazards.
Validation requires a scientific and technical review of the rationale behind each part of the HACCP plan (Slide 6). Validation activities may involve a scope, cost, and time commitment similar to the development of the original HACCP plan. In-plant validations should be performed before the HACCP plan is implemented, and when factors warrant. Validation activities can be performed by the HACCP team or by an individual qualified by training or experience.

**Validation involves establishing the scientific basis for the HACCP plan.**

Strategies that can be used to validate the HACCP plan include:

- using scientific principles and data,
- relying on expert opinion, or
- conducting in-plant observations or tests.

A number of different events or situations may affect when validation activities are needed (Slide 7). These factors could include changes in the raw materials, product or process; adverse review findings; recurring deviations; new scientific information about potential hazards or control measures; on-line observations; or new distribution or consumer-handling practices.

**Validation frequency:**
- Before the HACCP plan is implemented
- When factors warrant, such as:
  - changes in raw materials and/or suppliers
  - changes in product or process
  - adverse review findings
  - recurring deviations
  - new scientific information on hazards or control measures
  - on-line observations
  - new distribution or consumer handling practices

**Examples of Validation Procedures:**

- A seafood processor who is trying to determine an effective cook time and temperature to kill *Listeria monocytogenes* in their seafood products may use Table #A-3 in Appendix 4 of the Hazards Guide to validate their process. This table contains science-based information on the time and temperature combinations necessary for the destruction of *Listeria monocytogenes*, the primary pathogen of concern in the raw seafood products. The length of time at a particular internal product temperature that is needed to accomplish the recommended reduction in the number of *L. monocytogenes* is, in part, dependent upon the food in which it is being...
heated. The values in the table are generally conservative and apply to all foods. It may be possible to establish a shorter process time for a particular food by conducting scientific thermal death time studies, or by obtaining information from scientific studies that proves that the normal levels of this pathogen are lower than the expected levels.

- Processors may need to conduct in-plant thermal validation studies for a specific piece of equipment or process using science-based time-temperature scenarios to achieve the necessary reduction in pathogens. These in-plant validation studies should be done with the assistance of a food processing authority.
- A validation activity for a primary processor of molluscan shellfish could involve contacting the responsible authorities to identify the location of approved waters for raw shellfish before developing a harvest schedule. These authorities determine the status of harvesting waters using science-based sampling and testing protocols.
- A validation study for a brining process may involve science-based in-plant trials and product testing to determine the variables (e.g. size, fat content and amount of fish, salt level in the brine, brining time, etc.) that must be consistently achieved to assure that the required water phase salt will be reached in all finished products.

2) CCP Verification Activities

Several types of verification activities may be necessary for each CCP to ensure that the control procedures used are effective (Slide 8). A CCP verification procedure is needed to ensure that monitoring instruments are accurate and calibrated within appropriate ranges. CCP verification may also include targeted sampling and testing to demonstrate that the chosen critical limit is controlling the food safety hazard. Supervisory review of monitoring, corrective action and calibration or testing data is another type of CCP verification that is used to verify that the HACCP plan is working properly.

Slide 8

CCP verification activities:
- Calibration of process-monitoring devices
- Calibration record review
- Targeted sampling and testing
- CCP record review

Calibration of Process-monitoring Devices

Routine accuracy checks and periodic calibration of process-monitoring devices are CCP verification activities used to ensure that the measurements taken by the process-monitoring devices are accurate and reliable (Slide 9).

Accuracy checks and calibration are fundamental to the successful implementation and operation of the HACCP plan. If monitoring devices do not provide accurate measurements, then monitoring results will be unreliable. If
this happens, the CCP should be considered out of control (see Chapter 9 for corrective actions) since the last documented acceptable accuracy check and/or calibration.

For example, suppose a thermometer is used to measure the temperature of frying oil, which is part of the critical limit at a cooking CCP. The HACCP plan requires monthly calibration of this thermometer. If a scheduled calibration shows that the thermometer is actually reading 10°F less than the standardized thermometer, the accuracy of all monitoring results since the last monthly calibration is uncertain. Therefore, monitoring results for the last month are unreliable and it is impossible to know if the critical limit has been met during this period of time. Corrective action must be taken to determine if the products produced during this time period are safe and/or need to be recalled.

Calibration and accuracy are different but are related concepts (Slide 10). Ideally a measurement device is both accurate (correct or true) and precise (repeatable or reproducible). The accuracy and precision of a measurement process is usually established by measuring against a traceable reference standard.

Calibration involves determining, by measurement or comparison with two known standards, that the value of each reading on a particular measuring instrument is in fact correct or against a known calibrated instrument. For example, a thermometer could be calibrated by comparing it to a National Institute of Standards (NIST) traceable thermometer at two different temperatures in the range (above and below) in which it will be used.

Accuracy checks determine if the instrument is reading a true or correct value at a single point. For example, routine accuracy checks of a thermometer could involve immersing the probe into an ice slurry to determine if the thermometer measures a temperature of 32°F.

**Examples of accuracy checks and calibration activities:**

- A thermometer used to monitor temperature at a cooking CCP may be checked for accuracy by comparing it against a certified thermometer in a hot-water bath.
- The continuous temperature chart recorder on a pasteurizer may be compared during each batch against a certified accurate thermometer.
- A pH meter is calibrated against pH buffer standards of 7.0 and 4.0 when it is used to test products with a final pH of 4.2 to 4.8.

**Note**

In addition to the Hazards Guide, the National Seafood HACCP Alliance has prepared an online Compendium. This reference can be used to identify the types of equipment and methods that may be needed to develop effective monitoring and verification procedures.

**Web Link**

For a description of different types of thermometers to monitor refrigeration steps, go to: [http://seafood.ucdavis.edu/HACCP/Compendium/chapt06.htm](http://seafood.ucdavis.edu/HACCP/Compendium/chapt06.htm)

**Web Link**

For specific information about calibrating thermometers, go to: [http://seafood.ucdavis.edu/haccp/compendium/chapt02.htm#Thermometer%20calibration](http://seafood.ucdavis.edu/haccp/compendium/chapt02.htm#Thermometer%20calibration). There are also companies listed that offer thermometers for sale, including NIST thermometers.
Examples of accuracy checks:

- A thermometer used to routinely check the internal temperature of incoming fish is checked daily by immersing the probe in an ice slurry to determine that it reads 32°F.
- The accuracy of a metal detector is checked by running products with metal standards at the minimum detection limit through the unit.

**Frequency of Accuracy Checks and Calibration**

A number of factors should be considered when determining the frequency of accuracy checks and calibration for monitoring devices that is needed (Slide 11).
The design of a measuring instrument has to ensure that the device is capable of making accurate measurements when used within expected environmental conditions over some reasonable period of time.

Calibration frequency is dependent upon the type of device used, its condition and past performance, as well as the environment in which it will be used (operating environment). The reliability and sensitivity of the monitoring instrument should also be considered when determining the frequency of accuracy checks and calibration.

Consistent temperature variations away from the actual value (drift) found during checks and/or calibrations may indicate that more frequent calibration is needed or the device needs to be replaced (perhaps with a more durable device).

One of the most frequently used monitoring devices for perishable food products such as seafood is a thermometer. Some factors to consider when determining the frequency for thermometer accuracy checks and calibration include:

- **Inherent reliability**: Daily accuracy checks may be needed for the least reliable instruments (i.e. dial thermometers and bi-metallic). Periodic checks may be sufficient for more reliable instruments (i.e. digital thermometers with a history of good performance).
- **Manufacturer recommendation**: The design and expected conditions of use for each individual product is considered when manufacturers make accuracy and calibration recommendations. This information should be used to determine the frequency that is needed for these activities in the HACCP plan.

**Calibration Records and Review**

Records must be kept to document the results of accuracy checks and calibrations of monitoring devices (Slide 12). These records must be reviewed by a person who has the training or experience necessary to evaluate the results and determine that all monitoring instruments are accurate and properly calibrated. Reviewing these records may involve checking the dates and methods of calibration and the test results (e.g., equipment passing or failing). Calibration records must be reviewed within a reasonable time after completing the reports. These reviews are part of the plant’s verification activities. Inspectors and third-party auditors may also review calibration records.

**Web Link**

A Thermometer Calibration Guide and Poster are available from the following Web sites:


**Slide 12**

Calibration and accuracy checks records must:

1) Document results of accuracy checks and calibration procedures
2) Provide a reference to the standard
3) Be reviewed by qualified, trained personnel
Chapter 10

Targeted Sampling and Testing

Verification may also include targeted sampling, testing, and other periodic activities (Slide 13). For example, vendor compliance with a standard may be checked by targeted periodic sampling and testing when receipt of material is a CCP and purchase specifications outlined in a supplier guarantee are used as a critical limit. Typically, when a monitoring procedure does not involve a quantitative measurement, it should be coupled with a strong verification strategy. Similar to calibration records, sample test results must be reviewed within a reasonable time after completing the reports. These reviews are part of the plant’s verification activities.

Other examples of targeted sampling and testing:

• An initial, followed by quarterly sampling of shrimp at receiving and laboratory analysis for sulfite levels to verify that the incoming products do not exceed food safety specifications for this food additive.
• Periodic sampling and testing of water phase salt levels in finished smoked fish products to verify that the brining process is achieving the required water phase salt levels.
• Periodic sampling and histamine testing of fish from a harvest vessel to verify that the elements of the vessel’s handling and sampling procedures included in the processor’s critical limits do not create conditions that could lead to histamine formation.
• Periodic sampling and microbiological testing of cooked products to verify that the cooking process is sufficient to kill pathogens of concern.

CCP Record Review

The FDA regulation requires that all monitoring and corrective action records be reviewed within a week from the time that they are generated by a trained person (Slide 14). This review is a verification activity. These records are valuable tools that document that CCPs are operating within established safety parameters and that deviations are handled in a safe and appropriate manner. However, records alone are meaningless unless someone in a supervisory capacity reviews them on a periodic basis to “verify” that critical limits have been met and the HACCP plan is being followed.
3) HACCP Program Verification

In addition to the verification activities for CCPs, strategies should be developed for scheduled verification of the complete HACCP system (Slide 15). The frequency of a system-wide verification or reassessment shall be yearly, at a minimum, or whenever there is a system failure or a significant change in the product or process. The HACCP team is responsible for ensuring that this verification activity is performed. The HACCP team may contract with an independent third party to conduct system-wide verification activities.

**Slide 15**

HACCP system verification or reassessment frequency:
- Annually.
- Occurrence of a system failure or significant change in product or process.

System-wide verification activities may include on-site observations and record reviews (Slide 16). Reviews are usually performed by the HACCP Team or other unbiased individuals not responsible for performing the monitoring activities. It is recommended that the system verification occurs at a frequency that ensures the HACCP plan is being followed continuously. This frequency depends on a number of conditions, such as the variability of the process and product.

**Slide 16**

System-wide HACCP plan verification reviews include:
- Verifying that the hazard analysis and HACCP plan are still accurate, and
- Reviewing records to determine trends and verify that the plan is being followed.

Activities that should be conducted in a system-wide HACCP plan verification include:

- Check the accuracy of the product description and flow diagram.
- Check for new guidance or scientific information related to critical limits or other HACCP principles.
- Check that CCPs are monitored as required by the HACCP plan.
- Check that processes are operating within established critical limits.
- Check that appropriate corrective actions have been taken and verification activities have been completed.
- Check that records are completed accurately and at the time intervals required.
- Review sanitation records and other procedures that support the HACCP system.
- Review of consumer/customer complaints related to food safety.
During a system-wide HACCP plan verification, conduct a review of records to determine if the following have occurred:

- Monitoring activities have been performed at the locations specified in the HACCP plan.
- Monitoring activities have been performed at the frequencies specified in the HACCP plan.
- Corrective actions have been performed whenever monitoring indicated deviation from critical limits.
- Equipment has been calibrated at the frequencies specified in the HACCP plan.
- Check to be sure that all records are reviewed by a trained person within a week of the time they were generated.

It may be useful to include activities such as finished product testing and third-party audits when conducting a system wide HACCP verification (Slide 17).

**Slide 17**

Other system-wide verification strategies

- Finished product testing for microbiological, chemical or physical hazards
- Third-party audits

Finished product testing may include chemical or microbiological tests to ensure that the plan is controlling the food safety hazard identified at a CCP. As explained earlier, laboratory testing is generally ineffective for routine monitoring, but it can be used as a verification tool to determine if the overall operation is under control.

An independent third party audit can also be included in a system wide HACCP plan verification. Third party auditors can provide an unbiased assessment to help determine if the plan is working properly. Food processing authorities may also need to be consulted to re-validate a particular processing step.

Many different situations or conditions may trigger the need to reassess the HACCP plan to be sure that all food safety hazards are being effectively controlled (Slide 18).

**Slide 18**

Situations that may trigger a HACCP plan reassessment:

- A change in products or the process
- A change in the critical limit at a CCP
- Relocation of your plant
- Installation of a new piece of equipment
- A HACCP system failure
- Adverse findings from a regulatory inspection or third party audit
FDA’s Seafood HACCP regulation requires reassessment of the hazard analysis whenever there are any changes that could reasonably affect whether a food safety hazard now exists. The reassessment shall be performed by an individual or group of individuals who have been trained in HACCP principles.

4) Regulatory Verification

The major role of regulatory agencies in a HACCP program is to verify that HACCP plans are effective and are being followed and to make sure processors are in compliance with HACCP and other regulations (Slide 19). This inspection-based verification activity occurs on-site in the processing facility that has implemented the HACCP plan.

**Slide 19**

Regulatory agencies conduct inspection to verify that a processor:
- Has developed a HACCP plan that controls all significant food safety hazards;
- Has implemented the HACCP plan and it is consistently being used; and
- Is in compliance with HACCP and other regulations.

Verification procedures conducted by a regulatory agency may include:
- Review of the HACCP plan and any modifications
- Review of CCP monitoring records
- Review of corrective action records
- Review of the verification records
- Visual inspections of operations to determine if the HACCP plan is followed and records are properly maintained
- Random sample collection and analysis
- Evidence of training and that all training requirements have been met

HACCP plans are unique documents prepared by a processor to ensure the control of a specific process or procedure and to ensure compliance with regulations. The plans may contain proprietary information and must be appropriately protected by the regulatory agency. Agency personnel must have access to the HACCP plan plus monitoring, corrective action and verification records and other information pertinent to the HACCP plan that may be required for regulatory verification.

**Establishing a Schedule for Verification Activities**

Slide 20 provides an example of a company-established HACCP verification schedule. The frequency of all verification activities can change over time. A history of review findings that indicate that the processes are consistently in control may justify reducing the frequency. On the other hand, adverse findings, such as inconsistent monitoring activities, inconsistent record

**Note**

If you change/revise your HACCP plan, save the previous one for future reference.
keeping, or improper corrective actions may indicate that more frequent verification reviews are necessary and/or that the HACCP plan may need to be modified.

**Example of a company established HACCP verification schedule**

<table>
<thead>
<tr>
<th>Activity</th>
<th>Frequency</th>
<th>Responsibility</th>
<th>Reviewer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Verification activities scheduling</td>
<td>Yearly or upon HACCP program change</td>
<td>HACCP coordinator</td>
<td>Plant manager</td>
</tr>
<tr>
<td>Initial validation of HACCP plan</td>
<td>Prior to and during initial implementation of plan</td>
<td>Independent expert(s)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>HACCP team</td>
</tr>
<tr>
<td>Reassessment of HACCP plan</td>
<td>When critical limits changed, significant changes in process, equipment changes, after system failure, etc.</td>
<td>Independent expert(s)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>HACCP team</td>
</tr>
<tr>
<td>Verification of CCP monitoring as described in the plan (e.g., monitoring of patty cooking temperature)</td>
<td>According to HACCP plan (e.g., once per shift)</td>
<td>According to HACCP plan (e.g., line supervisor)</td>
<td>According to HACCP plan (e.g., quality control)</td>
</tr>
<tr>
<td>Review of monitoring, corrective action records to show compliance with the plan</td>
<td>Weekly</td>
<td>HACCP trained person</td>
<td>HACCP trained person</td>
</tr>
<tr>
<td>Comprehensive HACCP system verification</td>
<td>Yearly</td>
<td>HACCP team and/or independent expert(s)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>HACCP team</td>
</tr>
</tbody>
</table>

<sup>a</sup> May require additional technical expertise as well as laboratory and plant test studies.

**Examples of Verification Activities for Specific Hazards at a CCP**

Examples of appropriate verification procedures for various hazards at specific CCPs are provided in Slide 21. The verification procedures must be directly related to the critical limits, monitoring procedures and corrective action strategies identified in the HACCP plan.
Principle 6: Establish Verification Procedures

XYZ Seafood Company identified four critical controls points in the hazard analysis. There are three CCPs for the hazard of histamine: receiving, refrigerated storage and finished product storage. There is one CCP for the hazard of food allergens: weigh/pack/label. The HACCP team used the Hazards Guide to determine what verification procedures are needed in their HACCP plan.

<table>
<thead>
<tr>
<th>Significant Hazard</th>
<th>Critical Control Point</th>
<th>Critical Limits</th>
<th>Verification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aquaculture drugs</td>
<td>Receiving (from farm)</td>
<td>Suppliers certificate on file (indicating proper drug use).</td>
<td>Visit new suppliers within a year and existing suppliers on a pre-determined schedule to review drug use policies; and conduct quarterly sampling of raw material and test for drug residue likely to be present; and all records will be reviewed by a HACCP trained person once per week.</td>
</tr>
<tr>
<td>Natural toxins</td>
<td>Receiving (from harvester)</td>
<td>All shellstock tagged with the date and place of harvest, type and quantity of shellfish, and name or registration number of harvest vessel; and all shellstock from waters approved by State Shellfish Authority; and all shellstock from a licensed harvester.</td>
<td>Review all monitoring and corrective action records once per week.</td>
</tr>
<tr>
<td>Histamine</td>
<td>Receiving (from supplier)</td>
<td>Fish are completely surrounded by ice.</td>
<td>Check the accuracy of new thermometers before they are used and daily thereafter and calibrate thermometers once per year; and check internal temperature of iced fish at receipt before accepting fish from new suppliers and quarterly for existing suppliers to verify adequacy of ice; and all records will be reviewed by a trained person once per week.</td>
</tr>
<tr>
<td>C. botulinum toxin formation (in finished product)</td>
<td>Hot smoking</td>
<td>Internal fish temperature held at or above 145°F for at least 30 minutes.</td>
<td>Check the accuracy of the smokehouse temperature sensor before it is used and daily thereafter and calibrate at least once per year; and all records will be reviewed by a trained person once per week.</td>
</tr>
<tr>
<td>Pathogen-growth</td>
<td>Cooler storage</td>
<td>Cooler temperature not to exceed 40°F.</td>
<td>Check the accuracy of the cooler temperature sensor before it is used and daily thereafter and calibrate at least once per year; and all records will be reviewed by a trained person once per week.</td>
</tr>
</tbody>
</table>

Example: Fresh Mahi-mahi/XYZ Seafood Co.
Receiving CCP: The HACCP Team identified “Control Strategy 3, Transit Control,” in Chapter 7 (Histamine) of the Hazards Guide as the best control strategy for their situation. This strategy recommends verification procedures that include:

• Weekly review of Receiving Log (Monitoring record) and Corrective Action records
• Check internal temperature of fish at delivery for each new supplier and quarterly thereafter to ensure that icing procedures used by suppliers maintain product temperature
• Check accuracy of thermometer daily
• Calibrate thermometer used to check internal temperature annually

The verification procedures are entered in the HACCP plan form.

Refrigerated Storage and Finished Product Storage CCPs: The HACCP Team identified Storage Controls in Chapter 7 (Histamine) of the Hazards Guide as the best control strategy for both of their refrigerated storage CCPs. This strategy recommends verification procedures that include:

• Weekly review of the cooler ice log (monitoring record) and corrective action records
• Check internal temperature of fish quarterly to ensure that procedures used to ice fish for cooler storage maintains product temperature
• Check the accuracy of the thermometer for checking internal temperatures each time it is used
• Calibrate the thermometer used to check internal temperatures annually

The verification procedures are entered in the HACCP plan form.

Weigh/Pack/Label CCP: The HACCP Team identified Finished Product Labeling Controls in Chapter 19 (Major Food Allergens and Food Additives) of the Hazards Guide as the best control strategy for their situation. This strategy recommends verification procedures that include:

• Weekly review of Packing Room Log (monitoring record) and corrective action records

The verification procedures are entered in the HACCP plan form.

Example: Fresh Mahi-mahi/XYZ Seafood Co.

Complete HACCP Plan Form – Verification

Slide 22 – HACCP plan form for XYZ Seafood Company completed through Verification.
<table>
<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring</th>
<th>Corrective Action</th>
<th>Verification</th>
<th>Records</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receiving</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice at receipt.</td>
<td>Adequacy of ice surrounding mahi-mahi fillets at delivery</td>
<td>Visual check of adequacy of ice in a representative number of containers in each delivery</td>
<td>Every Delivery</td>
<td>Receiving Manager</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>If: the amount of ice is not adequate; Then: reject product, and call supplier to let them know CL was not met and provide product delivery specifications, and discontinue use of supplier until their transport procedures are corrected.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Weekly review of Cooler Ice Log (Monitoring Record) and Corrective action and Verification records</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Check internal temperature of fish quarterly to ensure that ice maintains product temperature</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Daily thermometer accuracy check.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Annual calibration of thermometer used to check internal temp.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Refrigerated Storage

<table>
<thead>
<tr>
<th>Refrigerated Storage</th>
<th>Histamine</th>
<th>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</th>
<th>Adequacy of ice surrounding mahi-mahi fillets</th>
<th>Visual check of adequacy of ice in a representative number of containers in cooler storage</th>
<th>At the beginning and end of the work day</th>
<th>Cooler Manager</th>
<th>Weekly review of Cooler Ice Log (Monitoring Record) and Corrective action and Verification records</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>If: the amount of ice is not adequate; Then: chill and hold the product until it can be evaluated based on its total time and temperature exposure, including exposures during prior processing operations, and add ice and make adjustments to the ice application process.</td>
<td></td>
<td></td>
<td>Weekly review of Cooler Ice Log (Monitoring Record) and Corrective action and Verification records</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Weekly review of Cooler Ice Log (Monitoring Record) and Corrective action and Verification records</td>
<td></td>
<td></td>
<td>Check internal temperature of fish quarterly to ensure that ice maintains product temperature</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Daily thermometer accuracy check.</td>
<td></td>
<td></td>
<td>Annual calibration of thermometer used to check internal temp.</td>
</tr>
<tr>
<td>Critical Control Point (CCP)</td>
<td>Significant Hazard(s)</td>
<td>Critical Limits for each Control Measure</td>
<td>Monitoring</td>
<td>Corrective Action</td>
<td>Verification</td>
<td>Records</td>
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<td>---------</td>
<td></td>
</tr>
<tr>
<td>Weigh/Pack/Label</td>
<td>Food Allergens</td>
<td>All finished product containers will be labeled with the correct market name of the fish.</td>
<td>The market name on each container of finished product</td>
<td>Visual check of a representative number of containers and their label</td>
<td>Each customer order</td>
<td>Packing Manager</td>
<td>Weekly review of Packing Room Log (Monitoring record) and Corrective action and Verification records</td>
</tr>
<tr>
<td>Finished Product Refrigerated Storage</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td>Adequacy of ice surrounding mahi-mahi fillets</td>
<td>Visual check of representative number of containers in cooler storage</td>
<td>At the beginning and end of the work day</td>
<td>Cooler Manager</td>
<td>Weekly review of Cooler Ice Log (Monitoring record), Corrective action and Verification records Check internal temperature of fish quarterly to ensure that ice maintains product temperature Daily thermometer accuracy check Annual calibration of thermometer used to check internal temp.</td>
</tr>
</tbody>
</table>

**Firm Name:** XYZ Seafood Company  
**Product:** Fresh mahi-mahi fillets  
**Firm Address:**  
238 Coastal Lane, Happy Beach, XX  
**Method of Storage and Distribution:** Stored and distributed buried in ice  
**Intended Use and Consumer:** To be cooked and consumed by the general public  
**Signature:**  
**Print name:**  
**Date:**
Chapter 11

Principle 7: Record-Keeping Procedures

Accurate recordkeeping is an essential part of a successful HACCP system (Slide 1). Recordkeeping is the seventh principle of HACCP.

In this chapter you will learn:
- What records are needed
- How to develop appropriate records
- How to conduct a record review
- How computerized records may be used

Types of Records Needed

Written records provide documentation of the HACCP plan, and demonstrate that critical limits have been met and appropriate corrective actions and verification procedures have been taken (Slide 2). Each type of record that is needed in a complete HACCP system is described, along with examples, in this chapter.

Six types of records are needed in a HACCP system:
1) The HACCP plan and supporting documentation
2) CCP monitoring records
3) Corrective action records
4) Verification records
5) Sanitation control records
6) Importer verification records
The FDA Seafood HACCP regulation requires firms to retain HACCP records for one year for fresh products and for two years for frozen or processed products with extended shelf-life. This regulation also requires that all records, plans, and procedures specified in a firm’s HACCP plan shall be made available for review and copying by public health officials at reasonable times. These requirements will be discussed in detail in Chapter 12.

1) The HACCP Plan and Supporting Documents

The written HACCP plan is a required record that describes exactly how all significant food safety hazards will be controlled (Slide 3). The HACCP plan is an official document that must include the name of the firm and its location, the products covered by the plan, and their method of storage, packaging, distribution, and intended use. The HACCP plan must also be signed and dated by a high level official or the most responsible individual to indicate that it has been accepted by the firm.

Documents that support the control strategies outlined in the HACCP plan could include any information used in completing the hazard analysis or in establishing critical limits (Slide 4).

Examples of HACCP Plan Support Documents:
- Data from published scientific studies
- Data from in-plant studies conducted by processing authorities
- Data from equipment manufacturers or other authorities
- Data gathered in the Preliminary Steps
- Pre-requisite programs including sanitation control procedures
- Written hazard analysis worksheets

Examples of documents that support the critical limits in a HACCP plan could include:
- Data from published scientific studies that demonstrate the adequacy of the control measures used to inhibit bacterial pathogen growth,
- Data from studies conducted by a processing authority to establish the safe shelf life of the product (if age of the product can affect safety),
- Data from an equipment manufacturer and an in-plant study conducted by a processing authority to establish the critical limit variables necessary for a cooking process to destroy bacterial pathogens.
- Data from an in-plant study conducted to determine the critical limit variables necessary to ensure that a brining process consistently achieves the minimum water phase salt levels necessary to prevent *C. botulinum* toxin production.
Examples of documents that support the overall HACCP plan and its strategies to control food safety hazards could include:

- A list of the firm’s HACCP team and their individual responsibilities
- The information gathered in the preliminary steps used to develop the HACCP plan
- Prerequisite programs including GMPs and sanitation control procedures
- A copy of the firm’s written hazard analysis worksheets

2) HACCP Monitoring Records

HACCP monitoring records are used to document that food safety hazards have been properly controlled at each CCP (Slide 5). Monitoring records show that critical limits are being met, and if not met, when appropriate corrective actions are needed. These records must be reviewed with sufficient frequency to ensure that food safety hazards are being controlled as required by the HACCP plan and regulatory requirements. Monitoring records are also used by regulators to determine whether a firm is in compliance with its written HACCP plan.

Note
Optional information, such as the Critical Limits from the HACCP plan, may be helpful to be included in the CCP monitoring record forms for reference.

Monitoring records can be routinely used by an operator or manager to determine if a process is approaching its critical limit. This enables the operator to make process adjustments before the critical limit is exceeded, which can reduce or eliminate the labor and material costs associated with corrective actions.

All monitoring information should be recorded at the time the observation is made. Accurate recordkeeping ensures the firm is meeting the critical limits and provides documentation that food safety hazards are being controlled. False or inaccurate records filled out before the operation takes place or ones that are completed later are inappropriate for a HACCP system and may lead to legal actions if found to be fraudulent.

Each monitoring record must be designed to capture the information required to document that the critical limit has been met at the CCP (Slide 6). The record must have a title that corresponds to the title of the record written in the HACCP plan. The actual measurement or observation that is taken must also be written on the record along with the time and date that the measurement or observation was taken and the signature or initials of the person who made it. There are additional record requirements in the FDA Seafood HACCP regulation related to the firm and its location, product identification, and record reviews that will be described in Chapter 12.
Examples of Monitoring Records for Specific Hazards at a CCP

Monitoring records must be designed to capture the measurements or observations that are included in the critical limit. Examples of the relationship between a CCP, critical limit, and monitoring records could include (Slide 7):

- A "Drug Use Certificate" is required from the supplier of farm raised fish to comply with the receiving CCP in order to reduce the hazard of aquaculture drugs to an acceptable level.
- A “Shellfish Receiving Log” is the record that would be filled out at a receiving CCP for shellfish when the critical limit requires properly filled tags that reduce the hazard of natural toxins to an acceptable level.

### Table: Examples of Monitoring Records for Specific Hazards at a CCP

<table>
<thead>
<tr>
<th>Significant Hazard</th>
<th>Critical Control Point</th>
<th>Critical Limits</th>
<th>Monitoring Record</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aquaculture drugs</td>
<td>Receiving (from farm)</td>
<td>Suppliers certificate on file (indicating proper drug use)</td>
<td>Suppliers certificate on file</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(indicating proper drug use)</td>
</tr>
<tr>
<td>Natural toxins</td>
<td>Receiving (from harvester)</td>
<td>All shellstock tagged with the date and place of harvest, type and quantity of shellfish, and name or registration number of harvest vessel AND All shellstock from waters approved by State Shellfish Authority AND All shellstock from a licensed harvester</td>
<td>Shellfish receiving log</td>
</tr>
<tr>
<td>Histamine</td>
<td>Receiving</td>
<td>Fish are completed surrounded by ice</td>
<td>Histamine fish receiving log</td>
</tr>
<tr>
<td>C. botulinum toxin formation (in finished product)</td>
<td>Hot smoking</td>
<td>Internal fish temperature held at or above 145°F for at least 30 minutes</td>
<td>Smokehouse temperature record chart</td>
</tr>
<tr>
<td>Pathogen growth</td>
<td>Cooler storage</td>
<td>Cooler temperature not to exceed 40°F</td>
<td>Cooler storage log</td>
</tr>
</tbody>
</table>
• A “Histamine Fish Receiving Log” is the record that would be filled out at a receiving CCP for histamine producing fish when the critical limit for a secondary processor is that all fish will be adequately surrounded with ice.
• A “Smokehouse Temperature Recording Log” is the record at a hot smoking CCP that would be examined and filled when the critical limit includes a minimum product internal temperature and processing time that is a necessary barrier to eliminate pathogens.
• A “Cooler Temperature Log” is the record that would be filled out at a refrigerated storage CCP when the critical limit is a maximum cooler storage temperature to prevent pathogen growth.

3) Corrective Action Records

Corrective action records were discussed in Chapter 9 and are repeated here to emphasize their importance in a firm’s HACCP system.

A corrective action must be taken each time a critical limit is not met and a corrective action record must show what products were affected, how the safety of these products was evaluated, the disposition of the affected product, the cause of the deviation, and how it was fixed (Slide 8).

Corrective actions records must capture information similar to that required for monitoring records, including:

• Product identification (e.g., product description, amount of product on hold)
• Description of the critical limit deviation
• Corrective action(s) taken including the final disposition of product and actions taken to prevent recurrence
• Results of the evaluation or testing of product placed on hold, if necessary
• Name and signature of the person responsible for the corrective action(s)
• Name and signature of the person reviewing the corrective action(s) report

A sample corrective action record was provided in Chapter 9 and will be reviewed at the end of this chapter.

4) Verification Records

Verification records were discussed in Chapter 10 and are repeated here to emphasize their importance in a firm’s HACCP system (Slide 9).
Records of routine and periodic verification activities must be kept to demonstrate that the HACCP plan has been implemented properly, monitoring measurements or observations are accurate and reliable, and the HACCP system is working as intended. Different records may be needed to capture the verification information that is specified in the HACCP plan.

**Slide 10**

Verification Records document the results of:
- Accuracy checks and calibration of process-monitoring instruments
- Record reviews
- Laboratory test results
- In-plant studies or challenge tests
- Audits and inspections

Examples of records for routine verification activities might include:

- Logs that document the results obtained when routine checks are made of the accuracy of thermometers, pH meters or other instruments used to monitor critical limits;
- Record review signature line and date on monitoring records to document the weekly review of monitoring and corrective action records by a trained person.

Examples of records for validation or periodic verification activities might include:

- A report that describes modifications made to the HACCP plan because of a change in products, ingredients, formulations, processes, or packaging and distribution methods;
- Processor audit records verifying supplier compliance with guarantees or certifications;
- Results of microbiological, chemical or physical tests of raw materials, in-process or finished products, or the plant environment;
- Results of equipment-evaluation tests, heat penetration or temperature distribution studies for thermal processes;
- Results from third party audits or regulatory agency inspections.

**Computerized Records**

Electronic or computerized records are acceptable in a HACCP system as long as they are equivalent to paper records and electronic signatures are equivalent to traditional handwritten signatures (Slide 11). If a company plans to use computerized records, they should review Title 21 CFR Part 11 for guidance (http://www.fda.gov).
Electronic record systems are classified as either open or closed. A closed system is one in which system access is controlled by the persons who are responsible for the content of the electronic records (e.g., a firm's HACCP coordinator). An open system is one in which system access is not controlled by the persons who are responsible for the content of electronic records (e.g., a software provider).

Controls are necessary for both types of systems to ensure that records are authentic, accurate, and protected from unauthorized changes (Slide 12). If a firm intends to implement an electronic record-keeping system, factors that must be considered in the design and implementation of the system include:

- Electronic records must be authentic, accurate and protected from unauthorized changes.
- They must be reviewed by management with sufficient frequency to ensure the firm's HACCP plan is being followed.
- They must be available for review and copying by public health authorities, if necessary.

An effective electronic record keeping system must:
- Be authentic, accurate and protected;
- Provide accurate and complete copies of records;
- Protect records for later retrieval;
- Limit access to authorized individuals;
- Provide a secure record audit trail; and
- Be reviewed by HACCP trained individual.

If a firm decides to use a specific electronic or computerized record-keeping system, it should be validated just like any other process or piece of equipment in the processing plant.

Recent advances in electronic communications makes the use of portable electronic devices attractive to firms who seek to reduce the amount of paper records that must be kept in a HACCP system. Any system that is used must ensure that the electronic records are equivalent to paper records and the electronic signatures are equivalent to traditional handwritten signatures.
Examples of Monitoring Records

Processors must design a form for each record listed in their HACCP plan to capture the information that is necessary to document monitoring results which show that critical limits have been met, what corrective actions have been taken, and that the necessary verification procedures have been completed. Because conditions in each facility are different, there is no single form that will be appropriate for all operations.

The following examples are generic records that illustrate basic design that can be used in a HACCP program:

**Routine Monitoring Records**

Daily Cooker Temperature Log Form (Slide 13)

This form documents the periodic recording of time and temperature under normal operating conditions of a cooker.

Continuous Temperature Record with Periodic Monitoring (Slide 14)

This record is used to continuously monitor the operations of a refrigerated storage. This record must be complemented by periodic visual checks by the operator to assure compliance with the CCP.

Weigh/Pack/Label Log (Slide 15)

This form documents the proper use of labels that identify potential hazards associated with the product or the product ingredients. Firms should affix one copy of the current label used to each report.

**Note**

Periodic readings of the temperature recording chart must be made to ensure that critical limits are met. Temperature monitoring records should be signed and dated by the line operator on the production day(s).

---

**Slide 13**

**Daily Cooker Temperature Log (Monitoring Record)**

<table>
<thead>
<tr>
<th>Form Title: Daily Cooker Temperature Log</th>
</tr>
</thead>
<tbody>
<tr>
<td>Firm Name: Firm Location:</td>
</tr>
<tr>
<td>Critical Limits: ≥ 212°F for ≥ 3 minutes</td>
</tr>
<tr>
<td>Monitoring Activities:</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Line Number</th>
<th>Product Code</th>
<th>Cooker Temp (°F)</th>
<th>Cook Time (minutes)</th>
<th>Critical Limit Met (Yes/No)</th>
<th>Line Operator (Initials)</th>
</tr>
</thead>
<tbody>
<tr>
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</tbody>
</table>

Reviewer Signature: Date of Review:
### Slide 14

**Continuous Temperature Record with Periodic Monitoring.**

![Temperature Chart](image)

XYZ Seafood Company  
238 Coastal Lane  
Happy Beach, XX  
Temperature Chart (cooler #1)  
2/12 - 2/19/11  
Reviewed By:  
Review Date:

### Slide 15

**Figure 3. Weigh/Pack/Label Log (Monitoring Record)**

<table>
<thead>
<tr>
<th>Form Title: Weigh/Pack/Label Log</th>
</tr>
</thead>
<tbody>
<tr>
<td>Firm Name:</td>
</tr>
<tr>
<td>Firm Location:</td>
</tr>
<tr>
<td>Product Identification:</td>
</tr>
<tr>
<td>Critical Limits: Proper label - seafood product market name and ingredients.</td>
</tr>
<tr>
<td>Monitoring Activities:</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Lot Number</th>
<th>Label Applied (Yes/No)</th>
<th>Label Type (description)</th>
<th>Line Operator (Initials)</th>
</tr>
</thead>
</table>

Reviewer Signature:  
Date of Review:
Corrective Action Records

Corrective Action Report (Slide 16)

This form is used to document the action taken when a critical limit is exceeded.

Corrective Action Report (Corrective Action Record)

<table>
<thead>
<tr>
<th>Form Title: Corrective Action Report Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>Firm Name:</td>
</tr>
<tr>
<td>Product Description:</td>
</tr>
<tr>
<td>Date:</td>
</tr>
<tr>
<td>Date and time of process deviation:</td>
</tr>
<tr>
<td>Describe the process deviation and what happened to the product?</td>
</tr>
<tr>
<td>What action(s) was taken to restore order to the process?</td>
</tr>
<tr>
<td>Name and signature of person reporting deviation and responsible for taking the correction action:</td>
</tr>
<tr>
<td>Amount of product affected by the process deviation:</td>
</tr>
<tr>
<td>Evaluation of product involved by the process deviation:</td>
</tr>
<tr>
<td>Final disposition of the affected product:</td>
</tr>
<tr>
<td>Reviewer Signature:</td>
</tr>
</tbody>
</table>

Routine Verification Records

Daily Thermometer Accuracy Log (Slide 17)

This form documents the daily accuracy check of all thermometers used in the daily process monitoring operations.

Thermometer Calibration Log (Slide 18)

This form records the quarterly calibration check of thermometers.
### Slide 17

**Daily Thermometer Accuracy Log (Verification Record)**

<table>
<thead>
<tr>
<th>Form Title: Daily Thermometer Accuracy Log</th>
</tr>
</thead>
<tbody>
<tr>
<td>Firm Name: Firm Location:</td>
</tr>
<tr>
<td>Product Identification:</td>
</tr>
<tr>
<td>Verification: Check each thermometer daily for accuracy. Temperature must be ± 2°F from the standard.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Instrument Number</th>
<th>Boiling Water Check</th>
<th>Critical Limit Met (Yes/No)</th>
<th>Line Operator (Initials)</th>
</tr>
</thead>
<tbody>
<tr>
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</table>

Reviewer Signature: Date of Review:

### Slide 18

**Quarterly Thermometer Calibration Log (Verification Record)**

<table>
<thead>
<tr>
<th>Form Title: Thermometer Calibration Log</th>
</tr>
</thead>
<tbody>
<tr>
<td>Firm Name: Firm Location:</td>
</tr>
<tr>
<td>Product Identification:</td>
</tr>
<tr>
<td>Verification: Check each thermometer daily for accuracy. Temperature must be ± 2°F from the standard.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date of Calibration</th>
<th>Instrument Number(s)</th>
<th>Method of Calibration</th>
<th>Calibration Results</th>
<th>Critical Limit Met (Yes/No)</th>
<th>Line Operator (Initials)</th>
</tr>
</thead>
<tbody>
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</table>

Reviewer Signature: Date of Review:
Periodic Verification and Validation Records

Finished Product Microbiological Evaluation Report (Slide 19)

This report documents the results of finished product laboratory analyses for total plate count (TPC), coliform bacteria, *Escherichia coli*, *Staphylococcus aureus* and *Salmonella*.

Raw Product Chemical Evaluation Report (Slide 20)

This report documents the results of laboratory analysis for sulfites in finished products.

Annual Thermal Process Validation Report (Slide 21)

This letter and supporting documents confirm the critical limits are scientifically valid and will deliver sufficient heat to destroy target organisms of public health concern.

Annual Thermal Equipment Validation Report (Slide 22)

This letter and supporting documents confirm that the temperature throughout the cooker is at or above the critical limit when the equipment is operating properly.

Annual HACCP Plan Verification Report (Slide 23)

This report document indicates that a firm has performed its annual HACCP system verification and the current HACCP plan was signed and dated.

Employee Training Record (Slide 24)

This report documents employee training activities.

---

**Finished Product Microbiological Evaluation Report (Verification Record)**

<table>
<thead>
<tr>
<th>Date of Sampling</th>
<th>Line Number</th>
<th>Total Plate Count (cfu/g)</th>
<th>Total coliforms (MPN/g)</th>
<th><em>E. coli</em> (Pos./Neg.)</th>
<th><em>S. aureus</em> (Pos./Neg.)</th>
<th><em>Salmonella</em> (Pos./Neg.)</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
<td>Reviewer Signature:</td>
<td>Date of Review:</td>
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</tbody>
</table>

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Slide 20

Raw Product Chemical Evaluation Report (Verification Record)

<table>
<thead>
<tr>
<th>Form Title: Chemical Evaluation Report</th>
</tr>
</thead>
<tbody>
<tr>
<td>Firm Name:</td>
</tr>
<tr>
<td>Firm Location:</td>
</tr>
<tr>
<td>Product Identification:</td>
</tr>
<tr>
<td>Verification: Sulfites must be declared</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Line Number</th>
<th>Sulfites (ppm)</th>
<th>Line Operator (Initials)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

Reviewer Signature: Date of Review:

Slide 21

Annual Thermal Process Validation Report (Verification Record)

May 9, 2011

John J. Smith, President
ABC Shrimp Company
One Saltwater Lane
Seaside, FL 33333

Dear Mr. Smith,

Heat penetration tests have been completed for your “Ready-to-eat, peeled and deveined shrimp” processed in a continuous steam cooker at your facility on April 19, 2010 using a portable data logger and 12 thermocouple leads.

Observations were made of internal product temperatures for six shrimp from individual lots of large (3.5 to 5.0 shrimp per oz.), medium (5.0 to 9.0 shrimp per oz.) and small (9.0 to 17.0 shrimp per oz.) processed in the steam cooker during production runs at 212°F for three minutes.

The internal temperature of large shrimp exceeded 165°F; medium shrimp, 170°F and small shrimp, 180°F. The internal product temperatures noted during these tests exceed your firm’s HACCP critical limits of an internal temperature of 165°F for 40 seconds.

Our studies revealed that shrimp processed at 212°F for three minutes delivered an internal product temperature above 165°F for a minimum of 40 seconds. These temperatures are equivalent to a 6-D process for elimination of Listeria monocytogenes.

These data serves as your annual thermal process validation study. If parameters change, such as cooking temperature, time, shrimp size, shrimp volumes, then you should repeat the thermal process validation study to ensure an adequate cook is being achieved in your process.

Sincerely,

I.M. Helpful
Seafood Processing and Research Unit
Your Seafood Processing Authority
Chapter 11

Slide 22

Annual Thermal Equipment Validation Report (Verification Record)

May 9, 2011

John J. Smith, President
ABC Shrimp Company
One Saltwater Lane
Seaside, FL 33333

Dear Mr. Smith,

Temperature distribution tests were performed on April 19, 2011 on the steam cooker located at ABC Shrimp Company in Sunshine, Florida.

Data was collected from ten thermocouples and continuous temperature logger during three production runs. Test results indicate that temperature distribution profiles in your cooker ranged from 212 to 214°F.

These studies show that your steam cooker, when run properly, continues to operate as designed.

Sincerely,

I.M. Helpful
Seafood Processing and Research Unit
Your State University

Slide 23

Annual HACCP Plan Verification Report (Verification Record)

<table>
<thead>
<tr>
<th>Annual HACCP Plan Verification Checklist</th>
<th>Date Task Completed:</th>
<th>Signature of Person who Completed the Task</th>
</tr>
</thead>
<tbody>
<tr>
<td>List of HACCP Team with Individual Responsibilities Updated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>List of Seafood Products and Processes in Place at Facility</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Product Flow Diagrams Updated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hazard Analysis Updated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HACCP Plan Updated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Good Manufacturing Practice Plan Updated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sanitation Standard Operating Practices Plan Updated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HACCP Plan Implemented</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reviewer Signature:</td>
<td>Date of Annual Review:</td>
<td></td>
</tr>
</tbody>
</table>
**Employee Training Report (Pre-requisite Document)**

<table>
<thead>
<tr>
<th>Employee Training Course</th>
<th>Date Completed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basic seafood HACCP and Sanitation SOPs workshop provided by Your University Extension program, AFDO accredited.</td>
<td>July 6, 2010</td>
</tr>
<tr>
<td>Computer operation of the pasteurizer, Best Yet Steam Cooker Co., John Jones, customer representative, three days on-the-job training.</td>
<td>September 23, 2010</td>
</tr>
<tr>
<td>Sanitation in the processing plant, 4-hour course, state inspection service, update.</td>
<td>March 12, 2010</td>
</tr>
<tr>
<td>Good manufacturing practices (GMPs) online course through Cornell University Distance Learning Center</td>
<td>September 10, 2010</td>
</tr>
</tbody>
</table>

**Example: Fresh Mahi-mahi/XYZ Seafood Co.**

XYZ Seafood Company identified four critical controls points in the hazard analysis. There are three CCPs for the hazard of histamine: receiving, refrigerated storage and finished product storage. There is one CCP for the hazard of food allergens: weigh/pack/label. The HACCP team used the Hazards Guide to determine what verification procedures are needed in their HACCP plan.

**Receiving CCP:** The HACCP Team identified “Control Strategy 3, Transit Control,” in Chapter 7 of the Hazards Guide as the best control strategy for their situation. This strategy recommends that processors keep a record (Receiving log) that documents: the number of containers examined, the number of containers in each delivery, and the results of checks for adequacy of ice.

**Refrigerated Storage and Finished Product Storage CCPs:** The HACCP Team identified Storage Controls in Chapter 7 of the Hazards Guide as the best control strategy for both of their refrigerated storage CCPs. This strategy recommends that processors keep a record (Cooler ice log) that documents: the number of containers examined, the adequacy of ice in each, the approximate number of containers in the cooler.

**Weigh/Pack/Label CCP:** The HACCP Team identified Finished Product Labeling Controls in Chapter 19 of the Hazards Guide as the best control strategy for their situation. This strategy recommends that processors keep a record (packing room log) that documents: the results of finished product labeling checks.
Examples of the monitoring records for XYZ Seafood Company are provided to illustrate how the basic record format developed in our previous examples can be adapted for different CCPs and critical limits. Similar adaptations can be made for the other records that would be needed including: a corrective action record, a thermometer accuracy check record, and an annual HACCP plan verification record.

**XYZ Seafood Company Record Examples**

Histamine Fish Receiving Log (Slide 25)

This record captures the results of the routine evaluations for the adequacy of ice surrounding all mahi-mahi fillets when they are received.

Histamine Fish Refrigeration Log (Slide 26)

This record documents the results of the routine evaluations for the adequacy of ice surrounding all mahi-mahi fillets that are stored in the refrigerated cooler.

Histamine Fish Packing Room Log (Slide 27)

This record captures the results of the routine monitoring of packaged product to ensure that it is properly labeled by market name for food allergen control.

---

**Histamine Fish Receiving Log (CCP Monitoring Record)**

<table>
<thead>
<tr>
<th>Date Received</th>
<th>Time</th>
<th>Number of containers received</th>
<th>Number of containers checked</th>
<th>Number of checked containers with adequate ice</th>
<th>Critical Limit Met? (Yes/No)</th>
<th>Receiving Manager (Initials)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2/11/11</td>
<td>8:05 AM</td>
<td>100</td>
<td>10</td>
<td>10</td>
<td>Y</td>
<td>p</td>
</tr>
<tr>
<td>2/12/11</td>
<td>7:30 AM</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>Y</td>
<td>p</td>
</tr>
<tr>
<td>2/13/11</td>
<td>11:00 AM</td>
<td>50</td>
<td>10</td>
<td>9</td>
<td>N</td>
<td>p</td>
</tr>
</tbody>
</table>

**Reviewer Signature:** [Signature]

**Date of Review:** 2/15/11
### Slide 26

**Histamine Fish Refrigeration Log (CCP Monitoring Record)**

<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Number of containers in cooler</th>
<th>Number of containers checked</th>
<th>Number of containers with adequate ice</th>
<th>Critical Limit Met? (Yes/No)</th>
<th>Receiving Manager (Initials)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</tr>
</tbody>
</table>

**Reviewer Signature:**

**Date of Review:**

### Slide 27

**Packing Room Log (CCP Monitoring Record)**

<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Number of containers in the order</th>
<th>Number of containers checked</th>
<th>Number of containers correctly labeled</th>
<th>Critical Limit Met? (Yes/No)</th>
<th>Packing Manager (Initials)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

**Reviewer Signature:**

**Date of Review:**
Example: Fresh Mahi-mahi/XYZ Seafood Co.

XYZ Seafood Company identified four critical controls points in the hazard analysis and has completed all elements of their HACCP plan except records. The final column of the HACCP plan form is completed using the information described above to describe the records that will be used to document their CCP monitoring procedures.

Complete HACCP Plan Form – Records

Slide 28 HACCP plan form for XYZ Seafood Company completed through Records
<table>
<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring</th>
<th>Corrective Action</th>
<th>Verification</th>
<th>Records</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receiving</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice at receipt.</td>
<td>Adequacy of ice surrounding mahi-mahi fillets at delivery</td>
<td>Visual check of adequacy of ice in a representative number of containers in each delivery</td>
<td>Every Delivery</td>
<td>Receiving Manager</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>If: the amount of ice is not adequate; <strong>Then:</strong> reject product and call supplier to let them know CL was not met and provide product delivery specifications, and discontinue use of supplier until their transport procedures are corrected.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Refrigerated Storage</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td>Adequacy of ice surrounding mahi-mahi fillets</td>
<td>Visual check of adequacy of ice in a representative number of containers in cooler storage</td>
<td>At the beginning and end of the work day</td>
<td>Cooler Manager</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>If: the amount of ice is not adequate; <strong>Then:</strong> chill and hold the product until it can be evaluated based on its total time and temperature exposure, including exposures during prior processing operations, and add ice and make adjustments to the ice application process.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Weekly review of Cooler Ice Log (Monitoring Record) and Corrective action and Verification records</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Receiving Log that documents: the number of containers examined; the number of containers in each delivery; and the results of checks for adequacy of ice.

Corrective Action records

Verification records

Cooler Ice Log that documents: the number of containers examined, the approximate number of containers in storage, and the results of checks for adequacy of ice.

Verification records
<table>
<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring</th>
<th>Corrective Action</th>
<th>Verification</th>
<th>Records</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weigh/Pack/Label</td>
<td>Food Allergens</td>
<td>All finished product containers will be labeled with the correct market name of the fish.</td>
<td>The market name on each container of finished product</td>
<td>Visual check of a representative number of containers and their label</td>
<td>Each customer order</td>
<td>Packing Manager</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Finished Product Refrigerated Storage</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td>Adequacy of ice surrounding mahi-mahi fillets</td>
<td>Visual check of representative number of containers in cooler storage</td>
<td>At the beginning and end of the work day</td>
<td>Cooler Manager</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Firm Name:** XYZ Seafood Company  
**Product:** Fresh mahi-mahi fillets  
**Firm Address:**  
238 Coastal Lane, Happy Beach, XX  
**Method of Storage and Distribution:** Stored and distributed buried in ice  
**Intended Use and Consumer:** To be cooked and consumed by the general public  
**Signature:** John Doe  
**Date:** 6/4/10  
**Print name:**
The Seafood HACCP Regulation

In December 1997, FDA’s regulation called “Procedures for the Safe and Sanitary Processing and Importing of Fish and Fishery Products” became final. The regulation is based on the seven principles of HACCP and is known as “the Seafood HACCP Regulation.” The Seafood HACCP Regulation will be referred to as “the regulation” for the rest of the chapter. A copy of the entire text of the regulation is found in Appendix 1 of this manual. This chapter will review the requirements of the regulation (Slide 1).

**Slide 1**

In this module, you will learn:
• The requirements of the regulation
• How to reference the specific requirements

**Regulation Format**

The regulation is part of Title 21 of the Code of Federal Regulations (CFR), Part 123, and is subdivided into three subparts and 13 sections. Subpart A is generally referred to as the “umbrella” section of the regulation as it applies to all processors of fish and fishery products. Subparts B and C are specific to processors of smoked fish and raw molluscan shellfish (Slide 2).

Subpart A contains twenty definitions that help a processor gain an understanding of the scope of the regulation and specific regulatory requirements (Slide 3).
Definitions 123.3

Of the twenty definitions, a few need to be emphasized (Slide 4).

Definitions 123.3

- certification number
- critical control point
- critical limit
- fish
- fishery product
- hazard
- importer
- molluscan shellfish
- preventive measure
- process-monitoring instrument
- processing
- processor
- scombroid toxin-forming species
- shall
- shellfish-control authority
- shellstock
- should
- shucked shellfish
- smoked or smoke-flavored fishery products
- tag

Regulatory terms “shall” and “should”
Regulations typically use the terms “shall” and “should”

“Shall” is used to state mandatory requirements of the regulations.
“Should” is used to state recommended or advisory procedures or to identify recommended equipment.

Two terms define which products are subject to the regulation (Slide 5).

**Slide 5**

Products that are subject to the regulation:
- Fish
- Fishery Product

**Fish** means freshwater or saltwater finfish, crustaceans, aquatic animal life (including alligators, frogs, aquatic turtles, jellyfish, sea cucumbers, sea urchins and roe) other than birds or mammals, and all mollusks, where such animal life is intended for human consumption. Note: it is important to carefully read the definition of fish and note that mollusks (molluscan shellfish) are considered fish for purposes of this regulation.

**Fishery product** means any human food product where fish is a characterizing ingredient, such as clam chowder or fish sauce.

Two terms define who must comply with the regulation (Slide 6).

**Slide 6**

Who must comply with the regulation:
- Importer 123.3 (g)
- Processor 123.3 (k) — domestic and foreign

An **Importer** means either the U.S. owner/consignee or the U.S. agent/representative of the foreign owner/consignee at the time of the product’s entry into the United States.

The person who is the owner or consignee at the time that the product is offered for entry is identified as the **importer** because: 1) that person has the ability to decide whether to offer the product for entry, and 2) that person is in a position to ensure that the product is processed under appropriate controls and to demonstrate this to FDA. The importer is responsible for ensuring that goods being offered for entry are in compliance with all laws affecting the importation. Ordinarily, the importer is not the custom-house broker, freight forwarder, carrier or steamship representative.

**Note**

This definition exempts products from the mandatory HACCP requirements that contain inconsequential amounts of fish. For example, Worcestershire sauce contains some anchovy paste but is not characterized by that ingredient.

**Note**

The ownership of an imported product can change many times in a short period of time after entry into the United States.
A **Processor** means any person engaged in commercial, custom or institutional **processing** of fish or fishery products **either in the United States or in a foreign country**.

One term defines what constitutes processing and is subject to the regulation (Slide 7).

**Slide 7**

What constitutes processing:
- Processing 123.3 (l)

**Processing** means handling, storing, preparing, heading, eviscerating, shucking, freezing, changing into different market forms, manufacturing, preserving, packing, labeling, dockside unloading, or holding fish or fishery products.

Processing is not defined by ownership of the product. A cold storage warehouse that stores product for different owners is responsible for complying with the regulation as they are “storing” fish and/or fishery products.

Certain processing practices/operations are exempt from the regulation (Slide 8).

- Fishing vessels and transporters who do not engage in processing are not subject to the regulation. However, primary processors that receive these products will need to evaluate the hazards associated with harvest and transportation and control significant hazards at receipt.
- Practices such as heading, gutting or freezing solely to prepare a fish for holding on board the harvest vessel are not subject to the regulation. For example, a fishing vessel may head and gut a halibut in order to better preserve it while holding on the vessel prior to unloading for further processing.
- Retail establishments are not subject to federal regulations, however, they must follow state and local government regulations. The Food Code (FDA's model food ordinance that many state and local regulatory authorities use in developing their food laws) requires that raw materials for retail establishments come from approved sources.

**Slide 8**

This regulation does not apply to:
- The harvest or transport of fish or fishery products
- Practices such as heading, eviscerating or freezing intended solely to prepare a fish for holding on a harvest vessel
- The operation of a retail establishment

**Note**

The term process also includes firms involved in developing fish and fishery products exclusively for market use or consumer tests (such as R&D products) as that is considered processing.

**Note**

Products that do not move in interstate commerce are not subject to this federal regulation. Products are considered to have entered into interstate commerce if raw materials, ingredients, packaging, etc. have originated outside the state. Products that strictly move in “intrastate” commerce are subject to state requirements. Many states have adopted HACCP regulations similar to the FDA Seafood HACCP Regulation.

**Note**

Fishing vessels that engage in processing – a.k.a. factory trawlers or catcher processors – are subject to the regulation.

**Note**

Aquaculture facilities that process at the same site as harvesting are subject to the regulation.

**Note**

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**Note**

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**Note**

Aquaculture facilities that process at the same site as harvesting are subject to the regulation.
Current Good Manufacturing Practices (21 CFR 123.5)

FDAs 21 CFR Part 110 current Good Manufacturing Practices (cGMPs) regulation outlines the conditions and practices the food industry must follow for processing safe food under sanitary conditions (Slide 9). The regulation is broad and includes all foods, including fish and fishery products. The regulatory requirements of the regulation are the basis for determining whether the facilities, methods, practices and controls used to process these products are safe and whether the products have been processed under sanitary conditions.

The seafood HACCP regulation compliments the cGMPs by requiring seafood processors to monitor and document the results of monitoring for eight key areas of sanitation derived from the cGMPs.

HACCP Plan 123.6

Hazard Analysis 123.6(a)

The regulation requires that every processor perform a hazard analysis (Slide 10). There are two major steps in a hazard analysis:

1) Determine whether there are hazards that are reasonably likely to occur
2) Identify control measures to control the identified hazards

Processors must consider hazards that are introduced both within and outside the processing plant and must consider food safety hazards that occur before, during or after harvest or transport. This means if you are a primary processor, in addition to considering hazards within your control, you must consider all hazards associated with your product that may occur prior to receipt. A secondary processor is responsible for considering hazards that might occur in-transit as well as hazards occurring within their processing facility.

Note

In Part 123.6 (a), the regulation references the term “preventive measure”. After the publication of the regulation, the term “preventive measure” was superceded by the more current term “control measure”.

Current Good Manufacturing Practices:
- Regulations found in Title 21, Part 110 of the Code of Federal Regulations
- Proper practices for the safe and sanitary handling of all foods

Slide 9

Hazard Analysis 123.6(a)

Every processor shall conduct, or have conducted for it, a hazard analysis.
The regulation defines a hazard that is reasonably likely to occur as “one for which a prudent processor would establish controls because experience, illness data, scientific reports or other information provide a basis to conclude that there is a reasonable possibility that it (the hazard) will occur in the particular type of fish or fishery product being processed in the absence of those controls (Slide 11).”

Slide 11

Determining those hazards that are “reasonably likely to occur.” Those “for which a prudent processor would establish controls.”

This means a prudent processor would establish controls because there is a reasonable possibility that a hazard will occur in the absence of controls. To make this decision, examine:

- Experience,
- Illness data,
- Scientific reports, and
- Other information. A useful source of information is the FDA Hazards Guide. It provides tables that outline the potential species- and process-related hazards that should be considered during the hazard analysis.

Even though every processor needs to conduct a hazard analysis, the regulation does not require it to be written. This is because it is the end product of the hazard analysis – the HACCP plan and its implementation – that will be evaluated by the regulatory authority.

However, a written hazard analysis will help the processor remember the thought process used to identify hazards and their controls. This is useful when periodic plan reassessments, a requirement of the regulation, are conducted and when the plan is reviewed by regulators.

To assist the industry, hazard analysis worksheet templates that can be used can be found in Appendix 2 of this manual.

**HACCP Plan 123.6(b)**

If the hazard analysis identifies one or more food safety hazards that are reasonably likely to occur, the processor shall have and implement a HACCP plan. Each HACCP plan must be specific to the processing location and each type of fish and fishery product. However, fish and fishery products that have the same hazards, same controls, same critical control points and same critical limits can be grouped into one HACCP plan (Slide 12).
The Contents of the HACCP Plan 123.6(c)

If a HACCP plan is needed it must list (Slide 13):

- The food-safety hazards that are reasonably likely to occur. Food safety hazards include biological, chemical and physical hazards.
- The critical control points
- The critical limits that will ensure that the identified hazard(s) are controlled
- The monitoring procedures that will ensure that the critical limits are being met. The frequency of monitoring must also be included.
- The pre-determined corrective actions unless the corrective action outlined in the regulation (21 CFR 123.7(c)) will be used.
- The verification procedures that ensure the system is operating according to plan. The frequency of verification must also be included.
- The records that will record the result of monitoring. Records must provide actual values or observations noted during monitoring.

These HACCP plan requirements are the same as the seven principles of HACCP discussed previously in this training.
Chapter 12

Signing and Dating the HACCP Plan 123.6(d)

The regulation requires that the HACCP plan be signed by the most responsible individual at the processing facility or a higher level official. The signature signifies that the plan has been accepted for implementation by the firm. The person who signs the plan is responsible for ensuring its accuracy, effectiveness and implementation and can be held responsible under the Seafood HACCP rule (Slide 14).

The HACCP plan shall be signed and dated:
- By the most responsible individual at the processing facility or a higher level official.
  - Signed and dated:
    - Upon initial acceptance.
    - Upon any modification.*
    - At least annually.*

* This is a verification requirement.

The signature and date on the HACCP plan is also outlined in the regulation. It must be signed and dated upon initial acceptance, upon any modification, and at least annually when it is reassessed.

Low Acid Canned Foods and Acidified Foods 123.6(e)

Processors of acidified and low acid canned foods are required to have controls in place for Clostridium botulinum under 21 CFR Part 113 and Part 114 (Slide 15). Because of this, processors who must comply with the requirements of Title 21 CFR Part 113 or 114 (acidified and low-acid canned foods) do not need to address the hazard of Clostridium botulinum in their HACCP plans. Their HACCP plans do not need to include controls to prevent that hazard, but they must continue to comply with 21 CFR Part 113 or 114. Other hazards may be reasonably likely to occur in an acidified or low-acid canned fishery product (e.g., histamine in canned tuna), and these must be addressed in the HACCP plan as appropriate.

Sanitation Controls and the HACCP Plan 123.6(f)

FDA recognizes that sanitation controls may be troublesome to manage in a HACCP plan. It is often difficult to determine appropriate critical limits and corrective actions for sanitation controls, particularly those relating to personnel hygiene (e.g., hand washing) (Slide 16). For this reason, the regulation

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Note

HACCP plans are not pre-approved by FDA before they are implemented by the processor. HACCP plans should not be submitted to the agency for review. FDA reached this decision because:
- HACCP plans and HACCP plan implementation are evaluated on-site, a process best accomplished during inspections of processing facilities.
- FDA does not have sufficient resources to review HACCP plans from all domestic and foreign seafood processors in advance of HACCP implementation by processors.

Note

When a processor controls a hazard such as cross-contamination of a ready-to-eat food through sanitation controls, the controls must be adequate to control the hazard and the monitoring procedures must be frequent enough to reliably indicate that the hazard is being controlled. In this example, cross-contamination of ready-to-eat foods is controlled.
The Seafood HACCP Regulation

does not require that sanitation controls be included in the HACCP plan. However, sanitation controls that are not in the plan must be monitored according to sanitation provisions of the regulation. Sanitation requirements are discussed in 21 CFR Part 123.11.

Legal Basis 21 CFR 123.6(g)

FDA issued the HACCP regulation under various sections of the Food Drug and Cosmetic Act, including, most significantly, sections 402(a)(1) and (a)(4). Together these sections state that a food is adulterated if it bears or contains any poisonous or deleterious substance that may render the food injurious to health, or if it is been prepared, packed or held under unsanitary conditions whereby it may have been contaminated with filth, or whereby it may have been rendered injurious to health (Slide 17).

Corrective Action 21 CFR 123.7

The regulation requires that a corrective action take place whenever a critical limit deviation has occurred (Slide 18). A corrective action that meets the requirements of the regulation must be designed to ensure that:

• No product enters commerce that is either injurious to health or is otherwise adulterated as a result of the deviation.
• The cause of the deviation is corrected.

Corrective Action 123.7
Whenever a deviation from a critical limit occurs, a processor shall take corrective action.
This two-fold approach ensures that the corrective action is applied to the **product** affected by the critical limit deviation, AND to the root cause of the **process** failure.

Processors have a choice of either 1) developing a predetermined corrective action plan in advance as part of their HACCP plans or 2) following the alternate procedure for corrective actions provided in the regulation (Slide 19). When a processor develops a plan in advance, he/she follows the plan that is appropriate when the deviation occurs. These corrective action plans become part of their HACCP plans as previously described in section 123.6(c).

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**Slide 19**

Corrective Actions — Two Choices:
1) Predetermined
2) Alternate Procedure – outlined in the regulation
   - Segregate and hold product
   - Determine product acceptability
   - Apply corrective action to product and process
   - Reassess the HACCP plan

---

Unusual situations may arise that may not be addressed by a predetermined corrective action plan. In these cases, the alternate corrective action procedure outlined in 21 CFR Part 123.7 (c) must be followed.

The alternate corrective action procedures listed in the regulation are:

- Segregating and holding the affected product until:
  - It is determined whether or not the product is safe for distribution. This decision must be made by someone who has suitable training or experience. This training or experience must be adequate for the person to understand the public health consequences of the critical limit deviation.
  - Corrective action is taken, as necessary, to ensure no unsafe product enters commerce.
- Corrective action is taken, as necessary, to fix the problem that caused the deviation.
- A reassessment is performed to determine whether the HACCP plan needs to be modified to reduce the risk that the deviation will happen again and modify the HACCP plan as necessary. This assessment and determination must be made by someone who has met the training requirements covered in section 123.10.

All corrective actions must be documented in records that include actions taken to ensure that affected unsafe product was not entered into commerce and the cause of the deviation was corrected.
Verification 123.8

Every processor must verify that the HACCP plan is adequate to control food safety hazards that are reasonably likely to occur, and that the plan is being effectively implemented (Slide 20). Verification must include, at a minimum, reassessment of the HACCP plan, ongoing verification activities, and record reviews.

Slide 20

Every processor shall verify:
- That the HACCP plan is adequate to control the food-safety hazards that are reasonably likely to occur; and
- That the HACCP plan is implemented effectively.

The HACCP plan must be reassessed at least once per year and whenever any changes occur that could affect the hazard analysis or the HACCP plan in any way. This could include changes in:

- Raw materials or source of raw materials
- Product formulation
- Processing methods or systems
- Finished product distribution systems
- The intended use or consumers of the finished product

The purpose of the reassessment is to ensure that the HACCP plan continues to be adequate to control the food-safety hazards which are reasonably likely to occur. It must be performed by an individual who meets the training requirements described in 21 CFR, part 123.10. If a processor has no HACCP plan because no significant hazards were identified, then the hazard analysis must be reassessed whenever any changes occur that could affect the original hazard analysis.

The regulation requires ongoing verification activities in addition to periodic reassessment. These ongoing activities are in keeping with the HACCP principle that verification must ensure that the HACCP plan process controls are effectively implemented on an ongoing basis. Verification activities must be listed in the HACCP plan. One of the functions of verification is to ensure a company’s adherence to its written HACCP plan. It is essential that HACCP plan components, including verification activities, are followed as written.

Consumer complaints must be reviewed by the processor to determine whether they relate to problems at a CCP (Slide 21).

Calibration of process monitoring instruments and routine accuracy checks are essential to the continued effective performance of CCPs. In addition to written calibration and accuracy check procedures, companies must perform these procedures at frequency intervals appropriate for the equipment and
instruments used to ensure the process controls continue to function as designed. Calibration is a verification procedure that must be listed in the HACCP plan.

End product or in-process testing methods are an optional verification strategy. However, end product and in-process tests can provide the processor with invaluable information that can be used to corroborate the ongoing effectiveness and adequacy of the process controls.

**Note**

Any end product or in process tests that are listed in the HACCP plan must be made available for review by the FDA.

**Slide 21**

Ongoing verification:
- Review of consumer complaints
- Calibration of process-monitoring instruments
- Periodic end-product and in-process testing (processor’s option)

**Records 21 CFR 123.9**

Records required by the regulation are (Slide 22):

- HACCP plan(s)
- Monitoring records
- Corrective action records
- Verification records
- Sanitation control records
- Importer verification records

**Slide 22**

Records required by the regulation:
- HACCP plan(s)
- Monitoring records
- Corrective action records
- Verification records
- Sanitation control records
- Importer verification records

All records required by the regulation shall be made available for review and copying by the regulatory authority.

Records required by the regulation **must** contain certain information (Slide 23):
- Name and location of the processor or importer
- Date and time of the activity being recorded
- Signature or initials of the person making the record
- Identity of the product and production code where appropriate
- Be completed at the time of the activity
The Seafood HACCP Regulation

The regulation requires that processors review certain records as part of verification (Slide 24). The purpose of these reviews is to ensure that the records are complete and that the activities occurred in accordance with the processor’s written procedures. The records must be reviewed by someone who meets the training requirements described in 21 CFR part 123.10.

Slide 23

Required information on each record:
- Name and location of the processor or importer
- Date and time of the activity being recorded
- Signature or initials of the person making the record
- Identity of the product and the production code where appropriate

Slide 24

Review of records:
- CCP monitoring and corrective action records – within one week
- Verification an in-process, end-product testing records – timely manner

Monitoring and corrective action records must be reviewed within one week of the day that the record was made. Calibration and in-process or end-product testing records must be reviewed in a timely manner.

Sometimes the performance of a verification procedure will indicate a potential public health problem. When this happens, the processor must follow the corrective action procedures described in 21 CFR part 123.7.

Records required by the regulation must be retained at the processing facility or importers place of business in the U.S. for at least one year after the date they were prepared in the case of refrigerated products and for or at least two years after the date they were prepared in the case of frozen preserved or shelf-stable products (Slide 25).

Slide 25

Record retention:
- One year for refrigerated products
- Two years for frozen or preserved products

Records that relate to the adequacy of equipment or process controls must be maintained at the processing facility or the importer’s place of business in the U.S. for at least two years.

Note

It may be desirable to review records more frequently to ensure control of products that are distributed daily, such as fresh seafood.
If permanent storage at the processing facility is not practical (e.g., a remote processing site or a processing vessel), the records may be transferred to some other facility at the end of the season. But the records must be made available for official review within a reasonable timeframe by a regulatory agency.

FDA has concluded that records and plans should be protected to the extent possible to promote the implementation of HACCP across the seafood industry. The regulation generally states that HACCP plans and records which come into FDA’s possession will not be available for public disclosure unless they have been previously disclosed, they relate to an abandoned product, and that they no longer represent a trade secret.

**Training 21 CFR 123.10**

The regulation requires that certain functions be performed by an individual trained in HACCP principles as applied to fish and fishery product processing. Processors can use a trained employee or a trained third party to perform these functions (Slide 26). The jobs may be done by one person or by several as long as they have been properly trained. The regulation defines a “HACCP-trained individual” as one “who has successfully completed training in the application of HACCP principles to fish and fishery product processing that is at least equivalent to that received under a standardized curriculum recognized as adequate by the U.S. Food and Drug Administration or who is otherwise qualified through job experience to perform these functions. Job experience will qualify if it has provided knowledge at least equivalent to that provided through the standardized curriculum.” This course material, developed by the National Seafood HACCP Alliance, is the standardized curriculum that has been recognized by the FDA.

<table>
<thead>
<tr>
<th>Slide 26</th>
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</thead>
<tbody>
<tr>
<td><strong>The HACCP-trained individual shall:</strong></td>
</tr>
<tr>
<td>• Develop the HACCP plan.</td>
</tr>
<tr>
<td>• Reassess and modify the HACCP plan and hazard analysis.</td>
</tr>
<tr>
<td>• Review HACCP records.</td>
</tr>
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</table>

**Sanitation Control Procedures (SCP) 21 CFR 123.11**

Sanitation is a prerequisite program that is necessary for the effective implementation of HACCP. In writing the regulation, FDA concluded that the GMP regulations (21 CFR 110) had not proven fully effective in encouraging seafood processors to take full responsibility for ensuring that sanitation in their plants consistently met minimum standards. For these reasons, the regulation requires that processors take certain actions to control sanitation conditions and practices (Slide 27).

These actions must be taken even if a processor determines there is no need for a HACCP plan. The sanitation requirements of the regulation may be made part of the processor’s HACCP plan or may be managed separately. Some processors may choose to use a combination of these approaches.
The regulation encourages but does not require “written” sanitation control procedures. However, the regulation does require that processors monitor the sanitary conditions and practices in their facility as well as correct any deficiencies that were noted during that monitoring. The regulation also requires written records documenting the results of the monitoring.

The eight key areas of sanitation include those sections of current Good Manufacturing Practices, which, if not controlled, would affect the safety of food (Slide 28). The purpose of the monitoring is to ensure that the requirements of the cGMPs are met and the purpose of the recordkeeping is to positively document the results of the monitoring. Monitoring frequencies are not specified but must be sufficient to ensure that the current GMP requirements are met.

- Processors should have written SCPs.
- Processor shall monitor and document sanitation control procedures.
- Processors shall correct sanitation deficiencies in a timely manner.

When sanitation conditions and practices are not met, they must be corrected in a timely manner. Records must be kept of the monitoring and the corrections. These records are subject to the same requirements as the HACCP records, except records review (verification) as outlined in 21 CFR part 123.8.

**Imported Products 21 CFR 123.12**

It has always been the importer's responsibility to offer products for entry into this country that are not adulterated under U.S. law. FDA's surveillance system for imports has traditionally consisted of reviews of customs entry forms for fish and fishery products being offered for entry into the United States, sensory analyses in the field (wharf examinations) and sample collections for laboratory analysis of products awaiting entry, and automatic detention of products with a history of problems.
Under the Seafood HACCP Regulation, additional HACCP controls are required for imported fish and fishery products as well as for domestic products. The definition of processor explicitly includes those who process seafood in foreign countries. Under this section, the U.S. Importer of Record bears the responsibility for verifying that foreign processors fully meet Seafood HACCP Regulations. To do so, the regulation requires that importers take certain steps to verify that their foreign suppliers meet the requirements of the regulation.

Importers may meet their obligation in one of two ways (Slide 29). They may import fish and fishery products that are covered by memorandums of understanding (MOU) between the United States and a foreign country. In this case, they would not need to take any other action to meet the requirements of the regulation.

If the U.S. does not have an MOU with the country of origin, importers must have and must implement written importer verification procedures that will ensure that the fish and fishery products offered for import into the United States were processed in accordance with the requirements of the regulation.

Importer Verification Procedures encompass three basic requirements (Slide 30). Importers must:

1) Have and implement written verification procedures that confirm the importer has written product specifications and has taken an affirmative step(s) to ensure that fish and fishery products offered for each entry into the U.S. have been processed in accordance with the regulation.
2) Have written product specifications for each of the products they are importing. Product specifications are designed to address those characteristics of the product that would be useful in providing assurance that the product is not adulterated under section 402 of the Federal Food, Drug and Cosmetic Act. This section relates to contaminants that may render the food injurious to health and to insanitary processing conditions. For example, it may be appropriate for a specification for frozen tuna steaks to include a maximum limit for histamine of 50 ppm.

In the FDA Hazards Guide, Appendix 5 (FDA and EPA Safety Levels in Regulations and Guidance) is an important reference that lists safety levels which can be used for product specifications of various fish and fishery products.

The product specification should cover any biological, chemical or physical hazard that might be reasonably likely to occur in the product being imported.
3) Take an affirmative step. An affirmative step may include any of those listed in the regulation, or other such verification procedures that provide an equivalent level of assurance that the fish or fishery product met the HACCP and sanitation requirements of the regulation (Slide 31).

**Slide 30**

**Importer Verification Procedures**

Importers must have:
1) Written verification procedures
2) Product specifications
3) Affirmative steps

**Slide 31**

Affirmative steps may include any of the following:
- Obtain foreign processor’s HACCP and sanitation monitoring records for the lot being entered
- Obtain continuing or lot-by-lot certificate from competent third party
- Regularly inspect foreign processor
- Obtain foreign processor’s HACCP plan and written guarantee that regulation is being met
- Test the product and obtain written guarantee that regulation is being met
- Perform other verification procedures that provide the equivalent level of assurance

The importer must keep records in English that document that the affirmative steps have been performed. The records must describe the results of the steps. These records are subject to the records requirements described in section 123.9. Importers that also process fish or fishery products must also meet the HACCP and sanitation requirements of the regulation for their processing operations.

An importer may hire a competent third party to perform verification activities. However, the importer remains responsible for demonstrating to FDA that the requirements have been met.

If an importer does not provide evidence that all of the fish and fishery products offered for entry into the United States have been processed under conditions that comply with this part of the regulation, the imported product will appear to be adulterated and will be denied entry.

**Web Link**

Letter of guarantee model can be found: [http://seafood.ucdavis.edu/haccp/guarant.htm](http://seafood.ucdavis.edu/haccp/guarant.htm)

**Note**

If an importer decides to obtain a foreign processor’s HACCP plan as part of their affirmative step, they should assess the plan to assure that the significant hazards have been listed. FDA can and will cite the importer for a HACCP plan not listing the significant hazards.

If an importer decides to obtain a continuing or lot-by-lot certificate from a competent third party as part of their affirmative step, the importer should review the document to ensure that the product is properly listed and the certificate date has not expired.

If an importer decides to regularly inspect the foreign processor as part of their affirmative step, they should include an inspection report describing their findings during the inspection.
Smoked and Smoke-Flavored Fishery Products 21 CFR 123.15 and 123.16

Subpart B of the regulation is specific to smoked fish (Slide 32). Smoked fish has been linked to a few cases of botulism. Clostridium botulinum, the bacterium that causes botulism, is prevented from growing in properly smoked fish by a combination of barriers, including salt, smoke, nitrite and, in the case of hot-smoked fish, heat. Careful control of these parameters is necessary to ensure the safety of the finished product. Such controls must be included in the HACCP plans of these products, unless the product is preserved by the addition of acid or heat under the controls required by the acidified or low-acid canned food regulations (21 CFR 113 and 114).

Slide 32

Smoked and Smoke-Flavored Fishery Products
- HACCP plan must include controls for Clostridium botulinum
toxin formation for the shelf life of the product under normal and moderate abuse conditions.
- Where product is subject to 21 CFR 113 or 114, the HACCP plan need not include such controls.

It is important to note that if there are other significant hazards, they must be included in the HACCP plan.

Raw Molluscan Shellfish 21 CFR 123.20 and 123.28 and
Control of Communicable Diseases – Molluscan Shellfish 1240.60

Subpart C of the regulation is specific to raw molluscan shellfish. Two interrelated programs have provided the basis for regulation of molluscan shellfish products by State Shellfish Control Agencies: the Interstate Shellfish Sanitation Conference (ISSC), and the National Shellfish Sanitation Program (NSSP).

The ISSC is an organization of state shellfish control agencies, the shellfish industry, and Federal agencies. The primary goal of the ISSC is to promote the adoption of uniform standards, rules, regulations, and procedures by state shellfish control agencies. Participation in the ISSC is voluntary, but it is supported by state shellfish control officials, participating nations, the shellfish industry, FDA, and the National Marine Fisheries Service.

The NSSP is a voluntary, tripartite program composed of state officials, the shellfish industry, and Federal agencies. FDA coordinates and administers the NSSP. In each participating state, one or more regulatory agencies manage the sanitation programs for domestic and imported shellfish.

With the advent of the regulation, in addition to compliance with State regulations, processors handling molluscan shellfish products must also comply with the Federal provisions outlined in 123.20-123.28.
The largest number of reported illnesses from consumption of seafood is caused by raw molluscan shellfish (oysters, clams and mussels) (Slides 33-39). These hazards are primarily introduced before the molluscan shellfish are harvested. The risk of occurrence of these hazards is reduced by ensuring that the molluscan shellfish come from sanitary growing waters. In most cases, the sanitary quality of molluscan-shellfish growing waters is determined by a state or national agency called a shellfish-control authority.

The regulation provides very specific requirements for controlling the source of origin for raw molluscan shellfish. It is important to note, however, that other hazards may also be reasonably likely to occur in these products, and they must be identified in the HACCP plan.

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**Slide 33**

Raw Molluscan Shellfish 123.20
- HACCP plans must include a means for controlling the origin of the raw molluscan shellfish.
- Where processing includes a treatment that ensures the destruction of vegetative cells of microorganisms of public health concern, the HACCP plan need not include controls on sources of origin.

**Slide 34**

Raw Molluscan Shellfish 123.28
Processors shall only process molluscan shellfish from:
- Growing waters approved by a shellfish-control authority
- Federal growing waters not closed by an agency of the federal government

**Slide 35**

Raw Molluscan Shellfish 123.28
Shellstock Receiving:
- If source is a harvester, harvester must be in compliance with any license requirement.
- If source is another processor, processor must be certified by a shellfish-control authority.
- Containers of shellstock must be properly tagged.
Raw Molluscan Shellfish 1240.60 (b)
Required information on tag:
• Date and place shellfish were harvested (state and site)
• Type and quantity of shellfish
• Harvester identification number, name of harvester or name or registration number of harvester’s vessel

Raw Molluscan Shellfish 123.28
Records for shellstock receiving must document:
• Date of harvest
• Location of harvest by state and site
• Quantity and type of shellfish
• Date of receipt by the processor
• Name of harvester, name or registration number of the harvester’s vessel or harvester’s identification number

Raw Molluscan Shellfish 123.28
Shucked molluscan shellfish containers must bear a label that contains:
• Name of packer or repacker
• Address of packer or repacker
• Certification number of packer or repacker

Raw Molluscan Shellfish 1240.60 (c)
Records for shucked product must document:
• Date of receipt
• Quantity and type of shellfish
• Name and certification number of the packer or repacker
A successful HACCP plan controls hazards (biological, chemical and physical) that are reasonably likely to occur to ensure that the food produced is safe to eat. The first part of this chapter introduces numerous resources that can assist in developing and modifying a seafood HACCP plan. The second part of this chapter provides instruction on how to use the FDA Fish and Fisheries Products Hazards and Controls Guidance to conduct your hazard analysis and develop a HACCP plan (Slide 1).

In this module, you will learn:
- What sources of information exist to help you identify seafood safety hazards and establish control measures
- How to use the Fish and Fishery Products Hazards and Controls Guide to identify hazards and establish control measures

Sources of Information on Seafood Hazards and Control Measures

You will need to perform a hazard analysis to decide if hazards are reasonably likely to occur in your products. Also, control measures need to be devised that make sense for your operations. To conduct a hazard analysis and develop a HACCP plan, gather information from a variety of sources (Slide 2) and choose the information that best applies to your situation. Some of the most useful sources of information are described in this chapter.
Personnel

The Seafood Processor

You and your employees know your operation better than anyone. Experience is an excellent source of information. You may already have knowledge about hazards that can affect your product, and you may have already implemented suitable controls.

Trade Associations

Trade associations can also provide useful information. Trade journals often provide general information on potential hazards and controls. Articles on specific processes or products also can be useful. Some trade organizations provide services such as consulting, educational programs, and publications that can help identify hazards and control measures.

Consultants and Auditors

Select HACCP consultants, firms, and auditors with expertise in the seafood HACCP regulation and control of seafood hazards. Consultants may be helpful in developing and reviewing your HACCP plan, particularly if you are just starting a new company or need expertise beyond your company’s abilities in complying with the seafood HACCP regulation, sanitation, sampling, etc. Auditors that you hire will have HACCP recommendations with the report that they provide.

University Sea Grant and Cooperative Extension Program Specialists

Many, but not all, coastal universities have seafood specialists within the Sea Grant or Land Grant Cooperative Extension programs. These programs provide outreach education and technical assistance to industry. Extension specialists and agents can assist in identifying potential hazards and control measures, but their availability may be limited.

Government Seafood Inspectors

Federal, state and local inspectors who visit your plant may point out potential hazards. It is the processor’s responsibility to implement effective control measures and to find solutions. The best time to access seafood inspectors for HACCP information is when they instruct at HACCP training courses.
**Suppliers, Buyers and Laboratory Analysts**

Suppliers of ingredients, cleaning materials, processing equipment and packaging materials, and analytical laboratories can provide information on potential hazards and control measures. A buyer's specification may point to a hazard in one of your products. For example, a buyer may require a *Salmonella*-free product. It is important to note, however, that not all buyer's specifications relate to safety. Analysts at analytical laboratories are a good source of information in developing validation studies and sampling programs.

**Processing Authorities**

Processing authorities use scientific methods to determine the proper parameters (temperature, atmosphere, oxygen, pH, etc.) to prevent, eliminate or reduce pathogens to acceptable levels. Processing authorities are a key source in validating the adequacy of your process and in providing technical advice for developing a HACCP plan, and implementing corrective action procedures.

**Publications**

The best and most accessible seafood HACCP resources available to develop and modify a HACCP plan are the two books that are provided in the Seafood HACCP Alliance basic HACCP course – the HACCP training curriculum, and the FDA Hazards Guide (Slide 3).

- **Slide 3**
  - Publications
    - The HACCP training curriculum
    - FDA Hazards Guide

- Chapters in this training curriculum give steps for developing a HACCP plan using a generic model of domestic mahi-mahi to be consumed by the general public. The first twelve chapters cover prerequisite programs; sanitation standard operating procedure requirements, introduction to hazards (biological, chemical, and physical); seven basic principles of HACCP; and seafood HACCP regulation. Through the seven principles of HACCP chapters, each chapter ends with either completed portions of the "hazard assessment" or the "HACCP plan" forms for the generic mahi-mahi model. Other models are available online from the Seafood HACCP Alliance for Training and Education (http://seafood.ucdavis.edu/haccpalliance.html).  
- Chapter 4, Seafood Safety Hazards, introduces the hazards that are common in fish and fishery products. Information is also provided on how these hazards can be controlled.  
- **The FDA Hazards Guide**
  - The Hazards Guide contains all seafood hazards currently identified by the FDA. The Hazards Guide or other scientific documentation may be used to conduct a hazard analysis and develop a HACCP plan.

**Note**

Additional hard copies of this training curriculum and the Hazards Guide are available for purchase from the University of Florida, IFAS Extension Bookstore (http://ifasbooks.com, follow the HACCP link)

**Web Link**

FDA Hazards Guide
Internet Resources

Information on key seafood hazards and controls sources is available free via the Internet. It is possible that over time the Web address may change or the information may be removed.

• The Seafood HACCP Alliance Compendium of Fish and Fishery Product Processes, Hazards, and Controls provides information that clarifies and supplements HACCP information from the training curriculum and Hazards Guide. The Compendium includes sections on seafood processes and controls, plus biological, chemical and physical hazards and controls. It provides the seafood industry with information on documented seafood process parameters, federal guidelines and tolerances for seafood contaminants, bacterial-growth parameters and recommended hazard-control operations. The compendium can assist in developing effective HACCP plans by providing scientific information on food-safety hazards and controls. The Compendium contains the following headings:

**Seafood Processes and Controls**
Chapter 1: Acidified, Fermented, and Salted Fish and Fisheries Products
Chapter 2: Battered Fish and Fisheries Products
Chapter 3: Cooked Fish and Fishery Products
Chapter 4: Dried Fish and Fishery Products
Chapter 5: Pasteurized Fish and Fisheries Products
Chapter 6: Refrigerated Fish and Fisheries Products
Chapter 7: Smoked Fish and Fisheries Products
Chapter 8: Vacuum and Modified Atmosphere Packaged Fish and Fisheries Products

**Biological Hazards and Controls**
Chapter 9: Aerobic Plate Count
Chapter 10: *Bacillus cereus*
Chapter 11: *Campylobacter* spp.
Chapter 12: *Clostridium botulinum*
Chapter 13: *Clostridium perfringens*
Chapter 14: Coliforms, Fecal Coliforms, and *Escherichia coli*
Chapter 15: *Listeria monocytogenes*
Chapter 16: Parasites
Chapter 17: *Salmonella* spp.
Chapter 18: *Shigella* spp.
Chapter 19: *Staphylococcus aureus*
Chapter 20: *Vibrio* spp.
Chapter 21: *Yersinia enterocolitica*

Web Link

Seafood Compendium
http://seafood.ucdavis.edu/haccp/compendium/compend.htm
Resources for Preparing HACCP Plans

Chemical Hazards and Controls
Chapter 22: Aquaculture Drugs
Chapter 23: Environmental Chemical Contaminants and Pesticides
Chapter 24: Food and Color Additives
Chapter 25: Methyl Mercury
Chapter 26: Natural Toxins
Chapter 27: Scombrotoxin (Histamine) Formation

Physical Hazards and Controls
Chapter 28: Hard or Sharp Objects

The format for each compendium chapter includes information that is available in the following areas:

- Potential Food Safety Hazards
- Control Measures
- Guidelines (Federal, State)
- Process Establishment
- Critical Aspects of Process
- Growth
- Heat Resistance
- Analytical Procedures
- HACCP Plan Examples
- Commercial Test Products
- Processes
- Time Temperature Monitoring Tags
- References

- The US Food and Drug Administration’s Web site provides quick and easy access to industry guidance, bulletins for health professionals, consumer education materials and other documents and data from FDA’s centers and offices. Key FDA seafood Web resources include:
  - The 2008 FDA Seafood List
    Contains FDA market names which generally should be used to label seafood products.
  - A list of aquaculture approved drugs can be found online from the FDA’s Center for Veterinary Medicine. At that site, the Aquatic Animal Drug Approval Partnership Program also makes available for download a full-sized reference poster, “Approved Drugs for Use in Aquaculture.”
  - The Center for Food Safety and Applied Nutrition (CFSAN)
    Is a gateway to numerous food safety pages including seafood, HACCP, labeling, allergens, bioterrorism, etc.
  - Fish and Fisheries Products Hazards and Controls Guidance
    (also known as the “Guide,” or the “Hazards Guide”) is a key HACCP resource to aid seafood processors and inspectors.
  - FDA Compliance Policy Guides (CPGs) and Import Alerts provide information on FDA compliance policy and insight to HACCP controls and new or unusual problems affecting import products that need to be addressed.
  - FDA Field Offices provides addresses and contact information of FDA offices located throughout the USA.

Web Links

FDA
http://www.fda.gov

Aquaculture Drugs
http://www.fda.gov/AnimalVeterinary/DevelopmentApprovalProcess/Aquaculture/ucm132954.htm

Seafood List
http://www.fda.gov/Food/GuidanceComplianceRegulatoryInformation/GuidanceDocuments/Seafood/ucm113260.htm

CFSAN
http://www.cfsan.fda.gov

CPGs

Import Alerts
http://www.fda.gov/ForIndustry/ImportProgram/ImportAlerts/default.htm

FDA Field Offices
http://www.fda.gov/ICECI/Inspections/IOM/ucm124008.htm
Web Links

Import Program
http://www.fda.gov/ForIndustry/ImportProgram/default.htm

Bad Bug Book
http://www.fda.gov/Food/FoodSafety/FoodborneIllness/FoodborneIllnessFoodbornePathogensNaturalToxins/BadBugBook/default.htm

Web Links

Listeria Guidance
http://www.fda.gov/Food/GuidanceCompliance/RegulatoryInformation/GuidanceDocuments/FoodProcessingHACCP/ucm073110.htm

HACCP Q&As
http://www.fda.gov/Food/GuidanceCompliance/RegulatoryInformation/GuidanceDocuments/Seafood/ucm176892.htm

Shellfish Shippers
https://info1.cfsan.fda.gov/shellfish/sh/shellfis.cfm

Managing Food Safety
http://www.fda.gov/Food/FoodSafety/RetailFoodProtection/ManagingFoodSafetyHACCPPrinciples/Operators/default.htm

- FDA Import Program provides links and information for seafood importers.
- The FDA Foodborne Pathogenic Microorganisms and Natural Toxins Handbook (known as the “Bad Bug Book”) contains basic facts regarding foodborne pathogenic microorganisms and natural toxins. It combines information from the Food and Drug Administration, the Centers for Disease Control and Prevention, the USDA Food Safety Inspection Service, and the National Institutes of Health. The “Bad Bug Book” provides food safety information on: pathogenic bacteria, enterovirulent Escherichia coli group (EEC group), parasitic protozoa and worms, viruses, natural toxins and other pathogenic agents. If you want background information on biological hazards in foods, consult the “Bad Bug Book” which provides the following available information per organism or toxin:

- Nature of the organism
- Nature of acute disease
- Nature of disease
- Diagnosis of human illness
- Associated foods
- Relative frequency of disease
- Course of disease and complications
- Target populations
- Food analysis
- Selected outbreaks
- Education and background resources
- Molecular structural data

Each chapter also has links to federal databases (CDC/MMWR, NIH/PubMed and AGRICOLA) which contain research articles on the organism or toxin.

- Guidance for Industry: Control of Listeria monocytogenes in Refrigerated or Frozen Ready-to-Eat Foods; Draft Guidance provides comprehensive information to industry on how to control Listeria monocytogenes.
- HACCP Regulation for Fish and Fishery Products, Questions and Answers for Guidance to Facilitate the Implementation of a HACCP System in Seafood Processing addresses many typical questions on clarifying interpreting and implementing the rule.
- Interstate Certified Shellfish Shippers List the shippers listed have been certified by regulatory authorities in the United States, Canada, Chile, Korea, Mexico and New Zealand under the uniform sanitation requirements of the national shellfish program.
- Managing Food Safety: A Manual for the Voluntary Use of HACCP Principles for Operators of Food Service and Retail Establishments is designed to assist the operators of retail and food service establishments in producing safe food.
Resources for Preparing HACCP Plans

- **Regulatory Fish Encyclopedia** assists with species identification and species names.
- **Seafood HACCP** contains links pertaining to the FDA regulation and resources.
- **Seafood HACCP: Final Rule** – 60 FR 665095 – December 18, 1995 (Regulation Title: Procedures for the Safe and Sanitary Processing and Importing of Fish and Fishery Products).
- **Seafood Information and Resources** contains links to seafood pathogens and contaminants, HACCP, inspection, imports, exports, compliance, federal and state programs, and resources.
- **The Seafood List, FDA’s Guide to Acceptable Market Names for Seafood Sold in Interstate Commerce** defines the different categories of names found in The Seafood List and outlines principles that can be used to label seafood species sold in the United States (U.S.) with an appropriate, non-misleading statement of identity.

- **The National Advisory Committee on Microbiological Criteria for Foods (NACMCF)** provides advice and recommendations to the secretaries of the Department of Agriculture and the Department of Health and Human Services concerning the development of microbiological criteria used to evaluate the safety and wholesomeness of food, including criteria for microorganisms that indicate whether food has been processed using GMPs. The NACMCF “Hazard Analysis Critical Control Point Principles and Application Guidelines” give an excellent overview of the HACCP principles with examples.

- **In the National Oceanic and Atmospheric Administration Seafood Inspection Program HACCP Quality Management Program** firms are audited by NOAA auditors for compliance with regulatory requirements and quality specifications for seafood. The HACCP QMP program combines the elements of internationally recognized quality standards with a HACCP-based audit program designed to verify a firm has controlled food safety issues. Under this program, firms can more fully integrate safety and quality controls under one comprehensive plan.

- **The National Shellfish Sanitation Program – Program Guide for the Control of Molluscan Shellfish** 2007 and the Interstate Shellfish Sanitation Conference. The NSSP is a cooperative federal/state/industry program established in 1925 to ensure the safety of molluscan shellfish. The program is described in the National Shellfish Sanitation Program Manual of Operations, Parts I and II. Part I is entitled “Sanitation of Shellfish Growing Areas,” and Part II is entitled “Sanitation of the Harvesting, Processing and Distribution of Shellfish.” The manual provides details on HACCP plan development and critical limits to ensure a safe product.

- **Sea Grant Library** (Digital HACCP Library) is a depository containing a wealth of HACCP related Web documents that are authored by Sea Grant programs.

Generic forms for conducting a hazard analysis and developing or modifying a HACCP plan are available from:

- this HACCP training curriculum
- the FDA Hazards Guide

**Web Links**

- **Fish Encyclopedia**
  http://www.fda.gov/Food/FoodSafety/Product-SpecificInformation/Seafood/RegulatoryFishEncyclopediaRFE/default.htm

- **Seafood HACCP**
  http://www.fda.gov/Food/FoodSafety/HazardAnalysisCriticalControlPointsHACCP/SeafoodHACCP/default.htm

- **Final Rule**
  http://www.fda.gov/Food/FoodSafety/Product-SpecificInformation/Seafood/SeafoodHACCP/ucm111304.htm

- **Information and Resources**
  http://www.fda.gov/Food/FoodSafety/Product-SpecificInformation/Seafood/default.htm

- **NACMCF Principles and Application Guidelines**
  http://www.fda.gov/Food/FoodSafety/HazardAnalysisCriticalControlPointsHACCP/ucm114868.htm

- **NOAA Quality Management Program**
  http://seafood.nmfs.noaa.gov/HACCPProgReq.pdf

- **NSSSP Program Guide**
  http://www.fda.gov/Food/FoodSafety/Product-SpecificInformation/Seafood/FederalStatePrograms/NationalShellfishSanitationProgram/Default.htm

- **Sea Grant Library**
  http://nsgl.gso.uri.edu/libraries/haccp.html
Chapter 13

Web Links

Generic HACCP Forms and Plans
http://seafood.ucdavis.edu/seafoodhaccp.html#section2

SHA HACCP Model Plans
http://seafood.ucdavis.edu/haccpalliance.html

Specifications Model
http://seafood.ucdavis.edu/haccp/fda-epa.htm

Letter of Guarantee
http://seafood.ucdavis.edu/haccp/guarant.htm

Aqua NIC
http://aquanic.org

CDC
http://www.cdc.gov

Morbidity and Mortality Weekly Report
http://www.cdc.gov/mmwr

CODEX Standards

FAO Technical Paper
http://www.fao.org/docrep/006/y4743e/y4743e00.htm

- the Seafood Network Information Center Web site:
  - generic HACCP flow diagram, hazard analysis and HACCP plan worksheets
  - generic HACCP plans
  - importer product specifications model
  - letter of guarantee model form for importers

There are numerous generic model HACCP plans available. They are useful in suggesting biological, chemical and physical hazards that may occur in your product or process. Do not simply select a generic HACCP plan and use it without modification. The plans should be adapted to reflect the conditions in your operation. The FDA knows how to recognize a copied HACCP model plan!

You must first independently conduct a hazard analysis of your species, processing (including ingredients), and environmental (sanitation control procedures) operations (see Principles 1-7 from Hazard Analysis to Record-Keeping Procedures). Then you need to develop your HACCP plan that is based on your specific plant location. The generic models give you an idea of the kinds of species and processing hazards that may occur in your situation. However, your HACCP plan must be tailored to your product and company’s site.

Miscellaneous Agency Resources

- The Aquaculture Network Information Center (AquaNIC) is a gateway to electronic resources on aquaculture. AquaNIC is linked to other aquaculture databases on the Internet. Primary funding of AquaNIC is through the NOAA Sea Grant College Program with secondary support from USDA North Central Regional Aquaculture Center.

- The Centers for Disease Control and Prevention is responsible for characterizing risk factors and prevention strategies for diseases that impact on public health. In addition, the CDC assists local health agencies in epidemiologic investigations of foodborne illness outbreaks. Certain diseases are reported to the CDC by state epidemiologists. The Morbidity and Mortality Weekly Report contains summaries of the information.

- The Codex Alimentarius Commission is sponsored by the Food and Agriculture Organization and the World Health Organization of the United Nations. Its purpose is to facilitate international trade by establishing uniform food standards. The commission has developed many standards and guidelines, including “Codex Alimentarius and Recommended International Code of Practice for Fish and Fishery Products.”

- The Food and Agriculture Organization of the United Nations (FAO) publication, Assessment and Management of Seafood Safety and Quality (FAO FISHERIES TECHNICAL PAPER 444) by H.H. Huss, L. Ababouch and L. Gram provides information on hazards and controls used in HACCP.
• FoodSafety.gov is a gateway to government food safety information, including links to foodborne pathogens, industry assistance, and government agencies.

• The National Academy of Sciences (NAS) received its congressional charter in 1863, which established it as a private, nonprofit organization designated as an official advisor to the federal government on science and technology matters. Its members include experts from many disciplines, including scientists, engineers, doctors, lawyers and corporate executives. The NAS Seafood Safety (1991) publication provides a good source of information about seafood hazards. In 2007 “Seafood Choices: Balancing Benefits and Risks," was edited by Malden C. Nesheim and Ann L. Yaktine and printed by the Institute of Medicine of the National Academies as a sequel to “Seafood Safety” and gives the perspective of nutritional benefits and risks to potential toxicants in seafood.

• The U.S. Department of Agriculture (USDA) HACCP regulation applies to processed meat and poultry products. Editor’s Note: You will need to conduct a search for meat and poultry HACCP on this Web site. If your company produces a seafood product which includes meat or poultry, or if your company processes both seafood and meat or poultry products, you also must comply with USDA HACCP requirements which are managed under the USDA Food Safety and Inspection Service.

• The University of California Seafood HACCP Discussion List (Internet Mailing List) is hosted at the University of California, Davis by the California Sea Grant Extension Program seafood technology unit. The HACCP discussion list provides an open Internet forum for seafood technology information exchange. Topics focus on seafood HACCP, safety, quality, processing, species, regulations, training programs, conferences and more. Subscriptions are free and available to anyone with e-mail access. Over 1,000 seafood professionals (industry personnel, inspectors, extension specialists and trade associations) from over 55 countries are subscribed. Questions posted on the discussion list usually receive a reply within a few hours to days.

• The University of California Seafood Network Information Center (SeafoodNIC) is sponsored by the University of California, Davis Sea Grant Extension Program seafood technology unit and the Department of Food Science and Technology. The SeafoodNIC hosts the National Seafood HACCP Alliance Web site, plus seafood-related guidelines and regulations, sanitation information, organizations, publications, and technical meetings. SeafoodNIC is also a gateway to other seafood resources on the Internet.

Web Links

Food Safety
http://www.foodsafety.gov

NAS Seafood Safety Publication
http://www.nap.edu/openbook.php?isbn=0309043875

Seafood Choices
http://books.nap.edu/catalog.php?record_id=11762

USDA Seafood HACCP
http://www.usda.gov

HACCP Discussion List
http://seafood.ucdavis.edu/listserv/listinfo.htm

SeafoodNIC
http://seafood.ucdavis.edu

Generic HACCP Plans
http://seafood.ucdavis.edu/haccp/Plans.htm
How to use the Fish and Fisheries Products Hazards and Controls Guidance

You have been provided a copy of the latest edition of the guide, along with your other training materials. You should use it as a reference tool during the practical exercise. Using the Hazards Guide, the key to getting started is to follow the 18 steps outlined in chapter 2 to develop a HACCP plan. The initial steps, Steps 1-11, complete the hazard analysis using a recommended hazard analysis form or worksheet (Slide 4). If no hazards are identified in these initial 11 steps, you can stop after completing the hazard analysis form. If hazards (biological, chemical, or physical) are identified, you will need to complete the remaining steps that build the HACCP plan. The Hazards Guide chapters and appendices provide background information and references on currently known seafood hazards and strategies for their control.

Slide 4

How the Hazards Guide Steps Correspond to the Hazard Analysis Worksheet

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Firm Name:</td>
<td>Firm Name: Step 1: Chapter 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Firm Address:</td>
<td>Firm Address: Step 1: Chapter 2 Step 5: Chapter 5 (Flow Diagram)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Product Description:</td>
<td>Product Description: Step 2: Chapter 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(1) Processing Step</td>
<td>Steps 7 and 8: Chapter 2, Chapter 3, and Appendix 6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(2) List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process.</td>
<td>Steps 9 and 10: Chapters 4-21</td>
<td>Steps 9 and 10: Chapters 4-21</td>
<td>Steps 9 and 10: Chapters 4-21</td>
</tr>
<tr>
<td>(3) Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No)</td>
<td>Justify the decision that you made in column 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(4)</td>
<td>What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard? (Yes or No)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(5)</td>
<td>Is this step a Critical Control Point? (Yes or No)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The Hazards Guide is designed so that a processor or regulator can look up the fish species and finished-product form of interest and identify potential food safety hazards. It is structured around the same hazard-analysis worksheet and HACCP plan form (Slide 5) that has been used throughout this course. In this way, the user is led through a series of decisions, such as whether a potential hazard is a significant hazard; what is the proper CCP; what critical-limit monitoring programs, corrective-action procedures and verification procedures are appropriate; and what records are necessary.
How the Hazards Guide Steps Correspond to the HACCP Plan Form

<table>
<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring</th>
<th>Corrective Action</th>
<th>Verification</th>
<th>Records</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step 12: Record processing step from hazard analysis form column 1 where CCP was identified</td>
<td>Step 12: Record corresponding hazard(s) from column 2 of hazard analysis form where CCP was identified</td>
<td>Step 13: Chapters 4-21 (section &quot;Set Critical Limits&quot;)</td>
<td>Step 14: Chapters 4-21 (section &quot;Establish Monitoring Procedures&quot;)</td>
<td>Step 15: Chapters 4-21 (section &quot;Establish Corrective Action Procedures&quot;)</td>
<td>Step 17: Chapters 4-21 (section &quot;Establish Verification Procedures&quot;)</td>
<td>Step 16: Chapters 4-21 (section &quot;Establish a Record-Keeping System&quot;)</td>
</tr>
</tbody>
</table>

Firm Name: Step 1: Chapter 2

Product: Step 2: Chapter 2

Firm Address: Step 1: Chapter 2

Method of Storage and Distribution: Step 3: Chapter 2

Intended Use and Consumer: Step 4: Chapter 2

Signature: Step 18: Chapter 2

Print name: ________________________________

Date: Step 18: Chapter 2
The recommendations included in the Hazards Guide are not, for the most part, binding FDA requirements. Use of the Hazards Guide in developing HACCP plans is not mandatory. It provides useful guidance, but seafood processors and importers are free to choose other control measures that provide an equivalent level of safety assurance to those listed in the guide. There may also be circumstances where a hazard identified in the guide may not apply to a product or species because of conditions specific to the processor.

Food-safety hazards can be introduced to a product because of the nature of the product (e.g., the species) or because of the way it is processed. The Hazards Guide refers to the first type as species-related hazards. It refers to the second type as process-related hazards. The Hazards Guide is set up in a way that lets you look up the species of interest (among the more than 350 listed) in a table. The table lists the potential species-related hazards that FDA has reason to believe exist for each species. You can also find the finished product of interest in another table. This table lists the potential process-related hazards that FDA has reason to believe exist for each finished product form. Processors must control both types of hazards.

The Hazards Guide then provides information to help processors and regulators decide if these potential hazards are reasonably likely to occur in any given circumstance. It further provides information about how the hazard might be controlled. These control options are not intended to be all-inclusive. Rather they represent the mechanisms that FDA is aware of that should prove effective in eliminating or minimizing the risk of a hazard developing in a product. In particular, the Hazards Guide provides information about critical limits that may be appropriate in certain circumstances. In some cases, the suggested critical limits are derived from a processing authority or existing tolerances or action levels. In other cases, they are derived from a review by FDA of the scientific and technical literature, conducted for the specific purpose of assisting in the development and review of HACCP plans.
Appendix 1

FDA’s Seafood HACCP Regulation

Title 21 of the Code of Federal Regulation
Part 123 – Fish and Fishery Products

Subpart A — General Provisions

§ 123.3 Definitions

The definitions and interpretations of terms in section 201 of the Federal Food, Drug, and Cosmetic Act (the act) and in part 110 of this chapter are applicable to such terms when used in this part, except where they are herein redefined. The following definitions shall also apply:

(a) **Certification number** means a unique combination of letters and numbers assigned by a shellfish control authority to a molluscan shellfish processor.

(b) **Critical control point** means a point, step or procedure in a food process at which control can be applied, and a food-safety hazard can be prevented, eliminated, or reduced to acceptable levels.

(c) **Critical limit** means the maximum or minimum value to which a physical, biological or chemical parameter must be controlled at a critical control point to prevent, eliminate or reduce to an acceptable level the occurrence of the identified food-safety hazard.

(d) **Fish** means fresh or saltwater finfish, crustaceans, other forms of aquatic animal life (including, but not limited to alligators, frogs, aquatic turtles, jellyfishes, sea cucumbers, sea urchins and roe of such animals) other than birds or mammals, and all mollusks, where such animal life is intended for human consumption.
(e) **Fishery product** means any human food product in which fish is a characterizing ingredient.

(f) **Food-safety hazard** means any biological, chemical or physical property that may cause a food to be unsafe for human consumption.

(g) **Importer** means either the U.S. owner or consignee at the time of entry into the United States or the U.S. agent or representative of the foreign owner or consignee at the time of entry into the United States, who is responsible for ensuring that goods being offered for entry into the United States are in compliance with all laws affecting the importation. For the purposes of this definition, ordinarily the importer is not the custom-house broker, the freight forwarder, the carrier or the steamship representative.

(h) **Molluscan shellfish** means any edible species of fresh or frozen oysters, clams, mussels, scallops or edible portions of such species, except when the product consists entirely of the shucked adductor muscle.

(i) **Preventive measure** means physical, chemical or other factors that can be used to control an identified food safety hazard.

(j) **Process-monitoring instrument** means an instrument or device used to indicate conditions during processing at a critical control point.

(k) (1) **Processing means**, with respect to fish or fishery products: Handling, storing, preparing, heading, eviscerating, shucking, freezing, changing into different market forms, manufacturing, preserving, packing, labeling, dockside unloading or holding.

(2) The regulations in this part do not apply to:

   (i) Harvesting or transporting fish or fishery products, without otherwise engaging in processing.

   (ii) Practices such as heading, eviscerating or freezing intended solely to prepare a fish for holding on a harvest vessel.

   (iii) The operation of a retail establishment.

(l) **Processor** means any person engaged in commercial, custom or institutional processing of fish or fishery products either in the United States or in a foreign country. A processor includes any person engaged in the production of foods that are to be used in market or consumer tests.

(m) **Scombroid toxin-forming species** means tuna, bluefish, mahi mahi, and other species, whether or not in the family Scombridae, in which significant levels of histamine may be produced in the fish flesh by decarboxylation of free histidine as a result of exposure of the fish after capture to temperatures that permit the growth of mesophilic bacteria.

(n) **Shall** is used to state mandatory requirements.
(o) **Shellfish control authority** means a federal, state or foreign agency, or sovereign tribal government, legally responsible for the administration of a program that includes activities such as classification of molluscan shellfish growing areas, enforcement of molluscan shellfish harvesting controls, and certification of molluscan shellfish processors.

(p) **Shellstock** means raw, in-shell molluscan shellfish.

(q) **Should** is used to state recommended or advisory procedures or to identify recommended equipment.

(r) **Shucked shellfish** means molluscan shellfish that have one or both shells removed.

(s) **Smoked or smoke-flavored fishery products** means the finished food prepared by:
   (1) Treating fish with salt (sodium chloride), and
   (2) Subjecting it to the direct action of smoke from burning wood, sawdust or similar material and/or imparting to it the flavor of smoke by a means such as immersing it in a solution of wood smoke.

(t) **Tag** means a record of harvesting information attached to a container of shellstock by the harvester or processor.

§ 123.5 **Current Good Manufacturing Practice**

(a) Part 110 of this chapter applies in determining whether the facilities, methods, practices and controls used to process fish and fishery products are safe and whether these products have been processed under sanitary conditions.

(b) The purpose of this part is to set forth requirements specific to the processing of fish and fishery products.

§ 123.6 **Hazard Analysis and Hazard Analysis Critical Control Point (HACCP) Plan**

(a) **Hazard analysis.** Every processor shall conduct or have conducted a hazard analysis to determine whether there are food-safety hazards that are reasonably likely to occur for each kind of fish and fishery product processed by that processor and to identify the preventive measures that the processor can apply to control those hazards. Such food-safety hazards can be introduced both within and outside the processing plant environment, including food-safety hazards that can occur before, during and after harvest. A food-safety hazard that is reasonably likely to occur is one for which a prudent processor would establish controls because experience, illness data, scientific reports or other information provide a basis to conclude that there is a reasonable possibility that it will occur in the particular type of fish or fishery product being processed in the absence of those controls.
(b) The HACCP plan. Every processor shall have and implement a written HACCP plan whenever a hazard analysis reveals one or more food-safety hazards that are reasonably likely to occur as described in paragraph (a) of this section. A HACCP plan shall be specific to:

1. Each location where fish and fishery products are processed by that processor; and
2. Each kind of fish and fishery product processed by the processor. The plan may group kinds of fish and fishery products together or group kinds of production methods together if the food-safety hazards, critical control points, critical limits and procedures required to be identified and performed in paragraph (c) of this section are identical for all fish and fishery products so grouped or for all production methods so grouped.

(c) The contents of the HACCP plan. The HACCP plan shall, at a minimum:

1. List the food-safety hazards that are reasonably likely to occur as identified in accordance with paragraph (a) of this section, and that must be controlled for each fish and fishery product. Consideration should be given to whether any food-safety hazards are reasonably likely to occur as a result of the following:
   (i) Natural toxins;
   (ii) Microbiological contamination;
   (iii) Chemical contamination;
   (iv) Pesticides;
   (v) Drug residues;
   (vi) Decomposition in scombroid toxin-forming species or in any other species where a food-safety hazard has been associated with decomposition;
   (vii) Parasites, where the processor has knowledge or has reason to know that the parasite-containing fish or fishery product will be consumed without a process sufficient to kill the parasites, or where the processor represents, labels or intends for the product to be so consumed;
   (viii) Unapproved use of direct or indirect food or color additives; and
   (ix) Physical hazards;
2. List the critical control points for each of the identified food-safety hazards, including as appropriate:
   (i) Critical control points designed to control food-safety hazards that could be introduced in the processing plant environment; and
   (ii) Critical control points designed to control food-safety hazards introduced outside the processing plant environment, including food-safety hazards that occur before, during and after harvest;
3. List the critical limits that must be met at each of the critical control points;
4. List the procedures, and frequency thereof, that will be used to monitor each of the critical control points to ensure compliance with the critical limits;
5. Include any corrective action plans that have been developed in accordance with § 123.7(b), to be followed in response to deviations from critical limits at critical control points;
(6) List the verification procedures, and frequency thereof, that the processor will use in accordance with § 123.8(a);

(7) Provide for a record-keeping system that documents the monitoring of the critical control points. The records shall contain the actual values and observations obtained during monitoring.

(d) **Signing and dating the HACCP plan.**

(1) The HACCP plan shall be signed and dated, either by the most responsible individual onsite at the processing facility or by a higher level official of the processor. This signature shall signify that the HACCP plan has been accepted for implementation by the firm.

(2) The HACCP plan shall be dated and signed:
   (i) Upon initial acceptance
   (ii) Upon any modification and
   (iii) Upon verification of the plan in accordance with § 123.8(a)(1).

(e) **Product subject to other regulations.** For fish and fishery products that are subject to the requirements of part 113 or 114 of this chapter, the HACCP plan need not list the food-safety hazard associated with the formation of *Clostridium botulinum* toxin in the finished, hermetically sealed container, nor list the controls to prevent that food-safety hazard. A HACCP plan for such fish and fishery products shall address any other food-safety hazards that are reasonably likely to occur.

(f) **Sanitation.** Sanitation controls may be included in the HACCP plan. However, to the extent that they are monitored in accordance with § 123.11(b) they need not be included in the HACCP plan and vice versa.

(g) **Legal basis.** Failure of a processor to have and implement a HACCP plan that complies with this section whenever a HACCP plan is necessary, otherwise operate in accordance with the requirements of this part, shall render the fish or fishery products of that processor adulterated under section 402(a)(4) of the act. Whether a processor’s actions are consistent with ensuring the safety of food will be determined through an evaluation of the processors overall implementation of its HACCP plan, if one is required.

§ 123.7 Corrective Actions

(a) Whenever a deviation from a critical limit occurs, a processor shall take corrective action either by:
   (1) Following a corrective action plan that is appropriate for the particular deviation, or
   (2) Following the procedures in paragraph (c) of this section.

(b) Processors may develop written corrective action plans, which become part of their HACCP plans in accordance with § 123.6(c)(5), by which they predetermine the corrective actions that they will take whenever there is a deviation from a critical limit. A corrective action plan that is appropriate for a particular deviation is one that describes the steps to be taken and assigns responsibility for taking those steps, to ensure that:
Appendix 1

(1) No product enters commerce that is either injurious to health or is otherwise adulterated as a result of the deviation; and
(2) The cause of the deviation is corrected.

c) When a deviation from a critical limit occurs and the processor does not have a corrective-action plan that is appropriate for that deviation, the processor shall:
   (1) Segregate and hold the affected product, at least until the requirements of paragraphs (c)(2) and (c)(3) of this section are met;
   (2) Perform or obtain a review to determine the acceptability of the affected product for distribution. The review shall be performed by an individual or individuals who have adequate training or experience to perform such a review. Adequate training may or may not include training in accordance with §123.10;
   (3) Take corrective action, when necessary, with respect to the affected product to ensure that no product enters commerce that is either injurious to health or is otherwise adulterated as a result of the deviation;
   (4) Take corrective action, when necessary, to correct the cause of the deviation;
   (5) Perform or obtain timely reassessment by an individual or individuals who have been trained in accordance with §123.10, to determine whether the HACCP plan needs to be modified to reduce the risk of reoccurrence of the deviation, and modify the HACCP plan as necessary.

d) All corrective actions taken in accordance with this section shall be fully documented in records that are subject to verification in accordance with §123.8(a)(3)(ii) and the record-keeping requirements of §123.9.

§123.8 Verification

(a) Overall verification. Every processor shall verify that the HACCP plan is adequate to control food-safety hazards that are reasonably likely to occur and that the plan is being effectively implemented. Verification shall include, at a minimum:

(1) Reassessment of the HACCP plan. A reassessment of the adequacy of the HACCP plan whenever any changes occur that could affect the hazard analysis or alter the HACCP plan in any way or at least annually. Such changes may include changes in the following: raw materials or source of raw materials, product formulation, processing methods or systems, finished product distribution systems, or the intended use or consumers of the finished product. Reassessment shall be performed by an individual or individuals who have been trained in accordance with §123.10. The HACCP plan shall be modified immediately whenever a reassessment reveals that the plan is no longer adequate to fully meet the requirements of §123.6(c).
(2) **Ongoing verification activities.** Ongoing verification activities including:

(i) A review of any consumer complaints that have been received by the processor to determine whether they relate to the performance of critical control points or reveal the existence of unidentified critical control points;

(ii) The calibration of process-monitoring instruments; and,

(iii) At the option of the processor, the performing of periodic end-product or in-process testing.

(3) **Records review.** A review, including signing and dating, by an individual who has been trained in accordance with § 123.10, of the records that document:

(i) The monitoring of critical control points. The purpose of this review shall be, at a minimum, to ensure that the records are complete and to verify that they document values that are within the critical limits. This review shall occur within 1 week of the day that the records are made;

(ii) The taking of corrective actions. The purpose of this review shall be, at a minimum, to ensure that the records are complete and to verify that appropriate corrective actions were taken in accordance with § 123.7. This review shall occur within one week of the day that the records are made; and

(iii) The calibrating of any process control instruments used at critical control points and the performing of any periodic end-product or in-process testing that is part of the processor’s verification activities. The purpose of these reviews shall be, at a minimum, to ensure that the records are complete, and that these activities occurred in accordance with the processor’s written procedures. These reviews shall occur within a reasonable time after the records are made.

(b) **Corrective actions.** Processors shall immediately follow the procedures in § 123.7 whenever any verification procedure, including the review of a consumer complaint, reveals the need to take a corrective action.

(c) **Reassessment of the hazard analysis.** Whenever a processor does not have a HACCP plan because a hazard analysis has revealed no food safety hazards that are reasonably likely to occur, the processor shall reassess the adequacy of that hazard analysis whenever there are any changes that could reasonably affect whether a food-safety hazard now exists. Such changes may include, but are not limited to changes in: raw materials or source of raw materials, product formulation, processing methods or systems, finished product distribution systems, or the intended use or consumers of the finished product. The reassessment shall be performed by an individual or individuals who have been trained in accordance with § 123.10.
(d) **Recordkeeping.** The calibration of process-monitoring instruments, and the performing of any periodic end-product and in-process testing, in accordance with paragraphs (a)(2)(ii) through (iii) of this section shall be documented in records that are subject to the record-keeping requirements of § 123.9.

§ 123.9 Records

(a) **General requirements.** All records required by this part shall include:

1. The name and location of the processor or importer;
2. The date and time of the activity that the record reflects;
3. The signature or initials of the person performing the operation; and
4. Where appropriate, the identity of the product and the production code, if any. Processing and other information shall be entered on records at the time that it is observed.

(b) **Record retention.**

1. All records required by this part shall be retained at the processing facility or importer’s place of business in the United States for at least one year after the date they were prepared in the case of refrigerated products and for at least two years after the date they were prepared in the case of frozen, preserved or shelf-stable products.
2. Records that relate to the general adequacy of equipment or processes being used by a processor, including the results of scientific studies and evaluations, shall be retained at the processing facility or the importer’s place of business in the United States for at least two years after their applicability to the product being produced at the facility.
3. If the processing facility is closed for a prolonged period between seasonal packs, or if record storage capacity is limited on a processing vessel or at a remote processing site, the records may be transferred to some other reasonably accessible location at the end of the seasonal pack but shall be immediately returned for official review upon demand.

(c) **Official review.** All records required by this part and all plans and procedures required by this part shall be available for official review and copying at reasonable times.

(d) **Public disclosure.**

1. Subject to the limitations in paragraph (d)(2) of this section, all plans and records required by this part are not available for public disclosure unless they have been previously disclosed to the public as defined in § 20.81 of this chapter or they relate to a product or ingredient that has been abandoned and they no longer represent a trade secret or confidential commercial or financial information as defined in § 20.61 of this chapter.
2. However, these records and plans may be subject to disclosure to the extent that they are otherwise publicly available, or that disclosure could not reasonably be expected cause a competitive hardship, such as generic-type HACCP plans that reflect standard industry practices.
(e) **Tags.** Tags as defined in § 123.3(t) are not subject to the requirements of this section unless they are used to fulfill the requirements of § 123.28(c).

(f) **Records maintained on computers.** The maintenance of records on computers is acceptable, provided that appropriate controls are implemented to ensure the integrity of the electronic data and signatures.

**§ 123.10 Training**

At a minimum, the following functions shall be performed by an individual who has successfully completed training in the application of HACCP principles to fish and fishery product processing at least equivalent to that received under standardized curriculum recognized as adequate by the U.S. Food and Drug Administration or who is otherwise qualified through job experience to perform these functions. Job experience will qualify an individual to perform these functions if it has provided knowledge at least equivalent to that provided through the standardized curriculum.

(a) Developing a HACCP plan, which could include adapting a model or generic-type HACCP plan, that is appropriate for a specific processor, in order to meet the requirements of § 123.6(b);

(b) Reassessing and modifying the HACCP plan in accordance with the corrective action procedures specified in § 123.7(c)(5), the HACCP plan in accordance with the verification activities specified in § 123.8(a)(1), and the hazard analysis in accordance with the verification activities specified in § 123.8(c); and

(c) Performing the record review required by § 123.8(a)(3); The trained individual need not be an employee of the processor.

**§ 123.11 Sanitation Control Procedures**

(a) **Sanitation SOP.** Each processor should have and implement a written sanitation standard operating procedure (herein referred to as SSOP) or similar document that is specific to each location where fish and fishery products are produced. The SSOP should specify how the processor will meet those sanitation conditions and practices that are to be monitored in accordance with paragraph (b) of this section.

(b) **Sanitation monitoring.** Each processor shall monitor the conditions and practices during processing with sufficient frequency to ensure, at a minimum, conformance with those conditions and practices specified in part 110 of this chapter that are both appropriate to the plant and the food being processed and relate to the following:

1. Safety of the water that comes into contact with food or food-contact surfaces, or is used in the manufacture of ice;
2. Condition and cleanliness of food-contact surfaces, including utensils, gloves and outer garments;
(3) Prevention of cross-contamination from insanitary objects to food, food-packaging material and other food-contact surfaces, including utensils, gloves and outer garments, and from raw product to cooked product;
(4) Maintenance of hand washing, hand sanitizing and toilet facilities;
(5) Protection of food, food-packaging material and food contact surfaces from adulteration with lubricants, fuel, pesticides, cleaning compounds, sanitizing agents, condensate and other chemical, physical and biological contaminants;
(6) Proper labeling, storage and use of toxic compounds;
(7) Control of employee health conditions that could result in the microbiological contamination of food, food-packaging materials and food-contact surfaces; and
(8) Exclusion of pests from the food plant. The processor shall correct in a timely manner, those conditions and practices that are not met.

c) **Sanitation control records.** Each processor shall maintain sanitation control records that, at a minimum, document the monitoring and corrections prescribed by paragraph (b) of this section. These records are subject to the requirements of §123.9.

d) **Relationship to HACCP plan.** Sanitation controls may be included in the HACCP plan, required by §123.6(b). However, to the extent that they are monitored in accordance with paragraph (b) of this section, they need not be included in the HACCP plan and vice versa.

### § 123.12 Special Requirements for Imported Products

This section sets forth specific requirements for imported fish and fishery products.

(a) **Importer verification.** Every importer of fish or fishery products shall either:

1. Obtain the fish or fishery product from a country that has an active memorandum of understanding (MOU) or similar agreement with the Food and Drug Administration, that covers the fish or fishery product and documents the equivalency or compliance of the inspection system of the foreign country with the U.S. system, accurately reflects the current situation between the signing parties, and is functioning and enforceable in its entirety; or
2. Have and implement written verification procedures for ensuring that the fish and fishery products that they offer for import into the United States were processed in accordance with the requirements of this part. The procedures shall list at a minimum:
   1. Product specifications that are designed to ensure that the product is not adulterated under section 402 of the Federal Food, Drug and Cosmetic Act because it may be injurious to health or have been processed under insanitary conditions, and,
(ii) Affirmative steps that may include any of the following:
(A) Obtaining from the foreign processor the HACCP and sanitation monitoring records required by this part that relate to the specific lot of fish or fishery products being offered for import;
(B) Obtaining either a continuing or lot-by-lot certificate from an appropriate foreign government inspection authority or competent third party certifying that the imported fish or fishery product is or was processed in accordance with the requirements of this part;
(C) Regularly inspecting the foreign processor’s facilities to ensure that the imported fish or fishery product is being processed in accordance with the requirements of this part;
(D) Maintaining on file a copy, in English, of the foreign processor’s HACCP plan, and a written guarantee from the foreign processor that the imported fish or fishery product is processed in accordance with the requirements of the part;
(E) Periodically testing the imported fish or fishery product, and maintaining on file a copy, in English, of a written guarantee from the foreign processor that the imported fish or fishery product is processed in accordance with the requirements of this part or;
(F) Other such verification measures as appropriate that provide an equivalent level of assurance of compliance with the requirements of this part.

(b) **Competent third party.** An importer may hire a competent third party to assist with or perform any or all of the verification activities specified in paragraph (a)(2) of this section, including writing the importer’s verification procedures on the importer’s behalf.

(c) **Records.** The importer shall maintain records, in English, that document the performance and results of the affirmative steps specified in paragraph (a)(2)(ii) of this section. These records shall be subject to the applicable provisions of § 123.9.

(d) **Determination of compliance.** There must be evidence that all fish and fishery products offered for entry into the United States have been processed under conditions that comply with this part. If assurances do not exist that the imported fish or fishery product has been processed under conditions that are equivalent to those required of domestic processors under this part, the product will appear to be adulterated and will be denied entry.
Subpart B – Smoked and Smoke-Flavored Fishery Products

§ 123.15 General

This subpart augments subpart A of this part by setting forth specific requirements for processing smoked and smoke-flavored fishery products.

§ 123.16 Process Controls

In order to meet the requirements of subpart A of this part, processors of smoked and smoke-flavored fishery products, except those subject to the requirements of part 113 or 114 of this chapter, shall include in their HACCP plans how they are controlling the food-safety hazard associated with the formation of toxin by Clostridium botulinum for at least as long as the shelf life of the product under normal and moderate abuse conditions.

Subpart C – Raw Molluscan Shellfish

§ 123.20 General

This subpart augments subpart A of this part by setting forth specific requirements for processing fresh or frozen molluscan shellfish, where such processing does not include a treatment that ensures the destruction of vegetative cells of microorganisms of public health concern.

§ 123.28 Source Controls

(a) In order to meet the requirements of subpart A of this part as they apply to microbiological contamination, chemical contamination, natural toxins, and related food safety hazards, processors shall include in their HACCP plans how they are controlling the origin of the molluscan shellfish they process to ensure that the conditions of paragraphs (b), (c), and (d) of this section are met.

(b) Processors shall only process molluscan shellfish harvested from growing waters approved for harvesting by a shellfish control authority. In the case of molluscan shellfish harvested from U.S. federal waters, the requirements of this paragraph will be met so long as the shellfish have not been harvested from waters that have been closed to harvesting by an agency of the federal government.

(c) To meet the requirements of paragraph (b) of this section, processors who receive shellstock shall accept only shellstock from a harvester that is in compliance with such licensure requirements as may apply to the harvesting of molluscan shellfish or from a processor that is certified by a shellfish control authority, and that has a tag affixed to each container of shellstock. The tag shall bear, at a minimum, the information required in § 1240.60(b) of this chapter. In place of the tag, bulk shellstock shipments may be accompanied by a bill of lading or similar shipping document that contains the information required in § 1240.60(b) of this chapter. Processors shall maintain records that document that all shellstock have met the requirements of this section. These records shall document:
(1) The date of harvest;
(2) The location of harvest by state and site;
(3) The quantity and type of shellfish;
(4) The date of receipt by the processor; and
(5) The name of the harvester, the name or registration number of the harvester’s vessel, or an identification number issued to the harvester by the shellfish control authority.

(d) To meet the requirements of paragraph (b) of this section, processors who receive shucked molluscan shellfish shall accept only containers of shucked molluscan shellfish that bear a label that complies with § 1240.60(c) of this chapter. Processors shall maintain records that document that all shucked molluscan shellfish have met the requirements of this section. These records shall document:
(1) The date of receipt,
(2) The quantity and type of shellfish, and
(3) The name and certification number of the packer or repacker of the product.

Part 1240 – Control of Communicable Diseases

2. The authority citation for 21 CFR part 1240 continues to read as follows: AUTHORITY: Secs. 215, 311, 361, 368 of the Public Health Service Act (42 U.S.C. 216, 243, 264, 271).

3. Section 1240.3 is amended by revising paragraph (r), and by adding new paragraphs (s), (t), and (u) to read as follows:

§ 1240.3 General Definitions

(r) Molluscan shellfish. Any edible species of fresh or frozen oysters, clams, mussels, and scallops or edible portions thereof, except when the product consists entirely of the shucked adductor muscle.

(s) Certification number means a unique combination of letters and numbers assigned by a shellfish control authority to a molluscan shellfish processor.

(t) Shellfish control authority means a federal, state, or foreign agency, or sovereign tribal government, legally responsible for the administration of a program that includes activities such as classification of molluscan shellfish growing areas, enforcement of molluscan shellfish harvesting controls, and certification of molluscan shellfish processors.

(u) Tag means a record of harvesting information attached to a container of shellstock by the harvester or processor.
4. Section 1240.60 is amended by revising the section heading, by redesignating the existing text as paragraph (a) and adding the word “molluscan” before the word “shellfish” the two times that it appears, and by adding new paragraphs (b), (c), and (d) to read as follows:

§ 1240.60 Molluscan Shellfish

(b) All shellstock shall bear a tag that discloses the date and place they were harvested (by state and site), type and quantity of shellfish, and by whom they were harvested (i.e., the identification number assigned to the harvester by the shellfish control authority, where applicable or, if such identification numbers are not assigned, the name of the harvester or the name or registration number of the harvester’s vessel). In place of the tag, bulk shellstock shipments may be accompanied by a bill of lading or similar shipping document that contains the same information.

(c) All containers of shucked molluscan shellfish shall bear a label that identifies the name, address and certification number of the packer or repacker of the molluscan shellfish.

(d) Any molluscan shellfish without such a tag, shipping document, or label, or with a tag, shipping document or label that does not bear all the information required by paragraphs (b) and (c) of this section, shall be subject to seizure or refusal of entry, and destruction.
Appendix 2

HACCP Worksheets

Worksheets are recommended to document the hazard analysis and final HACCP plans. The hazard analysis should contain certain information to justify the identification of the proper food safety hazards and critical control points. Information in the HACCP plan must explain the details for each HACCP step. There is no standardized or mandated format for the worksheets, but the information should be arranged in a progressive manner that clearly explains the thought process for the hazard analysis and the individual steps in the HACCP plan.

The following worksheets are provided as recommended examples. The information is arranged in a similar manner, but the layouts are in either a landscape or portrait form to suit individual preferences.

**Special Note:** These recommended worksheets can be copied for routine use, but if they are used for official use they must include details that identify the commercial firm and related information. The additional information must include:

- Form title,
- Firm name and location,
- Time and dates,
- Product identification,
- Signature and date (HACCP plan).
Product Description Form for Fish and Shellfish Species

<table>
<thead>
<tr>
<th>Type of Seafood Product (Species name)</th>
<th>Where Product Is Purchased (Source)</th>
<th>How Product Is Received</th>
<th>How Product Is Stored</th>
<th>How Product Is Shipped</th>
<th>How Product is Packaged</th>
<th>Intended Use</th>
<th>Intended Consumer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>From Fisherman</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>Refrigerated</td>
<td>Air Packed</td>
<td>Raw, RTE</td>
<td>General Public</td>
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<td>From Fish Farm</td>
<td>Iced</td>
<td>Iced</td>
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<td>Reduced-Oxygen/</td>
<td>Cooked, RTE</td>
<td>At-risk Population</td>
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<td>From Processor</td>
<td>Frozen</td>
<td>Frozen</td>
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<td>Raw, to-be-cooked</td>
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## Hazard Analysis Worksheet – Traditional Method

<table>
<thead>
<tr>
<th>Processing Step</th>
<th>List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process.</th>
<th>Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No)</th>
<th>Justify the decision that you made in column 3</th>
<th>What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard?</th>
<th>Is this step a Critical Control Point? (Yes or No)</th>
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| Processing Step | List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process. | Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No) | Justify the decision that you made in column 3 | What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard? | Is this step a Critical Control Point? (Yes or No) | }

### Hazard Analysis Worksheet – Inclusive Method

**Firm Name:**

**Product Description:**

**Firm Address:**

**Method of Storage & Distribution:**

**Intended Use & Consumer:**
## Hazard Analysis Worksheet – Inclusive Method

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<td>(1) Ingredient/processing step</td>
<td>(2) Identify potential hazards introduced, controlled or enhanced at this step.</td>
<td>(3) Are any potential food-safety hazards significant? (Yes/No)</td>
<td>(4) Justify your decision for column 3</td>
<td>(5) What control measure(s) can be applied to prevent the significant hazards?</td>
<td>(6) Is this step a critical control point? (Yes/No)</td>
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# HACCP Plan Form

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**Firm Name:**

**Product:**

**Firm Address:**

**Method of Storage and Distribution:**

**Intended Use and Consumer:**

**Signature:** ____________________________ **Date:** ____________________________

**Print name:** ____________________________
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<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
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#### Monitoring Corrective Action Verification Records

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#### Receiving Histamine Mahi-mahi fillets

- **Refrigerated Storage**
  - Histamine Mahi-mahi fillets are completely surrounded with ice at receipt.

- **Finished Product**
  - Refrigerated Storage
  - Histamine Mahi-mahi fillets are completely surrounded with ice throughout storage time.

#### Weigh/Pack/Label Food Allergens

- All finished product containers will be labeled with the correct market name of the fish.
Current Good Manufacturing Practices (21CFR110)


Source: 51 FR 24475, June 19, 1986, unless otherwise noted.

Subpart A – General Provisions

§ 110.3 Definitions.

The definitions and interpretations of terms in section 201 of the Federal Food, Drug, and Cosmetic Act (the act) are applicable to such terms when used in this part. The following definitions shall also apply:

(a) **Acid foods or acidified foods** means foods that have an equilibrium pH of 4.6 or below.

(b) **Adequate** means that which is needed to accomplish the intended purpose in keeping with good public health practice.

(c) **Batter** means a semifluid substance, usually composed of flour and other ingredients, into which principal components of food are dipped or with which they are coated, or which may be used directly to form bakery foods.

(d) **Blanching** except for tree nuts and peanuts, means a prepackaging heat treatment of foodstuffs for a sufficient time and at a sufficient temperature to partially or completely inactivate the naturally occurring enzymes and to effect other physical or biochemical changes in the food.

(e) **Critical control point** means a point in a food process where there is a high probability that improper control may cause, allow, or contribute to a hazard or to filth in the final food or decomposition of the final food.
(f) **Food** means food as defined in section 201(f) of the act and includes raw materials and ingredients.

(g) **Food-contact surfaces** are those surfaces that contact human food and those surfaces from which drainage onto the food or onto surfaces that contact the food ordinarily occurs during the normal course of operations. “Food-contact surfaces” includes utensils and food-contact surfaces of equipment.

(h) **Lot** means the food produced during a period of time indicated by a specific code.

(i) **Microorganisms** means yeasts, molds, bacteria, and viruses and includes, but is not limited to, species having public health significance. The term “undesirable microorganisms” includes those microorganisms that are of public health significance, that subject food to decomposition, that indicate that food is contaminated with filth, or that otherwise may cause food to be adulterated within the meaning of the act. Occasionally in these regulations, FDA used the adjective “microbial” instead of using an adjectival phrase containing the word microorganism.

(j) **Pest** refers to any objectionable animals or insects including, but not limited to, birds, rodents, flies, and larvae.

(k) **Plant** means the building or facility or parts thereof, used for or in connection with the manufacturing, packaging, labeling, or holding of human food.

(l) **Quality control operation** means a planned and systematic procedure for taking all actions necessary to prevent food from being adulterated within the meaning of the act.

(m) **Rework** means clean, unadulterated food that has been removed from processing for reasons other than insanitary conditions or that has been successfully reconditioned by reprocessing and that is suitable for use as food.

(n) **Safe-moisture level** is a level of moisture low enough to prevent the growth of undesirable microorganisms in the finished product under the intended conditions of manufacturing, storage, and distribution. The maximum safe moisture level for a food is based on its water activity ($a_w$). An $a_w$ will be considered safe for a food if adequate data are available that demonstrate that the food at or below the given $a_w$ will not support the growth of undesirable microorganisms.

(o) **Sanitize** means to adequately treat food-contact surfaces by a process that is effective in destroying vegetative cells of microorganisms of public health significance, and in substantially reducing numbers of other undesirable microorganisms, but without adversely affecting the product or its safety for the consumer.

(p) “**Shall**” is used to state mandatory requirements.
“Should” is used to state recommended or advisory procedures or identify recommended equipment.

Water activity \( (a_w) \) is a measure of the free moisture in a food and is the quotient of the water vapor pressure of the substance divided by the vapor pressure of pure water at the same temperature.

§ 110.5 Current good manufacturing practice.

(a) The criteria and definitions in this part shall apply in determining whether a food is adulterated \((1)\) within the meaning of section 402(a)(3) of the act in that the food has been manufactured under such conditions that it is unfit for food; or \((2)\) within the meaning of section 402(a)(4) of the act in that the food has been prepared, packed, or held under insanitary conditions whereby it may have become contaminated with filth, or whereby it may have been rendered injurious to health. The criteria and definitions in this part also apply in determining whether a food is in violation of section 361 of the Public Health Service Act \((42 \text{ U.S.C. 264})\).

(b) Food covered by specific current good manufacturing practice regulations also is subject to the requirements of those regulations.

§ 110.10 Personnel.

The plant management shall take all reasonable measures and precautions to ensure the following:

(a) Disease control. Any person who, by medical examination or supervisory observation, is shown to have, or appears to have, an illness, open lesion, including boils, sores, or infected wounds, or any other abnormal source of microbial contamination by which there is a reasonable possibility of food, food-contact surfaces, or food-packaging materials becoming contaminated, shall be excluded from any operations which may be expected to result in such contamination until the condition is corrected. Personnel shall be instructed to report such health conditions to their supervisors.

(b) Cleanliness. All persons working in direct contact with food, food-contact surfaces, and food-packaging materials shall conform to hygienic practices while on duty to the extent necessary to protect against contamination of food. The methods for maintaining cleanliness include, but are not limited to:

1. Wearing outer garments suitable to the operation in a manner that protects against the contamination of food, food-contact surfaces, or food-packaging materials.
2. Maintaining adequate personal cleanliness.
3. Washing hands thoroughly (and sanitizing if necessary to protect against contamination with undesirable microorganisms) in an adequate hand-washing facility before starting work, after each absence from the work station, and at any other time when the hands may have become soiled or contaminated.
(4) Removing all unsecured jewelry and other objects that might fall into food, equipment, or containers, and removing hand jewelry that cannot be adequately sanitized during periods in which food is manipulated by hand. If such hand jewelry cannot be removed, it may be covered by material which can be maintained in an intact, clean, and sanitary condition and which effectively protects against the contamination by these objects of the food, food-contact surfaces, or food-packaging materials.

(5) Maintaining gloves, if they are used in food handling, in an intact, clean, and sanitary condition. The gloves should be of an impermeable material.

(6) Wearing, where appropriate, in an effective manner, hair nets, headbands, caps, beard covers, or other effective hair restraints.

(7) Storing clothing or other personal belongings in areas other than where food is exposed or where equipment or utensils are washed.

(8) Confining the following to areas other than where food may be exposed or where equipment or utensils are washed: eating food, chewing gum, drinking beverages, or using tobacco.

(9) Taking any other necessary precautions to protect against contamination of food, food-contact surfaces, or food-packaging materials with microorganisms or foreign substances including, but not limited to, perspiration, hair, cosmetics, tobacco, chemicals, and medicines applied to the skin.

(c) **Education and training.** Personnel responsible for identifying sanitation failures or food contamination should have a background of education or experience, or a combination thereof, to provide a level of competency necessary for production of clean and safe food. Food handlers and supervisors should receive appropriate training in proper food handling techniques and food-protection principles and should be informed of the danger of poor personal hygiene and insanitary practices.

(d) **Supervision.** Responsibility for assuring compliance by all personnel with all requirements of this part shall be clearly assigned to competent supervisory personnel.

[51 FR 24475, June 19, 1986, as amended at 54 FR 24892, June 12, 1989]

§ 110.19 Exclusions.

(a) The following operations are not subject to this part: Establishments engaged solely in the harvesting, storage, or distribution of one or more “raw agricultural commodities,” as defined in section 201(r) of the act, which are ordinarily cleaned, prepared, treated, or otherwise processed before being marketed to the consuming public.

(b) FDA, however, will issue special regulations if it is necessary to cover these excluded operations.
Subpart B – Buildings and Facilities

§ 110.20 Plant and grounds.

(a) Grounds. The grounds about a food plant under the control of the operator shall be kept in a condition that will protect against the contamination of food. The methods for adequate maintenance of grounds include, but are not limited to:

1. Properly storing equipment, removing litter and waste, and cutting weeds or grass within the immediate vicinity of the plant buildings or structures that may constitute an attractant, breeding place, or harborage for pests.
2. Maintaining roads, yards, and parking lots so that they do not constitute a source of contamination in areas where food is exposed.
3. Adequately draining areas that may contribute contamination to food by seepage, foot-borne filth, or providing a breeding place for pests.
4. Operating systems for waste treatment and disposal in an adequate manner so that they do not constitute a source of contamination in areas where food is exposed.

If the plant grounds are bordered by grounds not under the operator’s control and not maintained in the manner described in paragraph (a) (1) through (3) of this section, care shall be exercised in the plant by inspection, extermination, or other means to exclude pests, dirt, and filth that may be a source of food contamination.

(b) Plant construction and design. Plant buildings and structures shall be suitable in size, construction, and design to facilitate maintenance and sanitary operations for food-manufacturing purposes. The plant and facilities shall:

1. Provide sufficient space for such placement of equipment and storage of materials as is necessary for the maintenance of sanitary operations and the production of safe food.
2. Permit the taking of proper precautions to reduce the potential for contamination of food, food-contact surfaces, or food-packaging materials with microorganisms, chemicals, filth, or other extraneous material. The potential for contamination may be reduced by adequate food safety controls and operating practices or effective design, including the separation of operations in which contamination is likely to occur, by one or more of the following means: location, time, partition, air flow, enclosed systems, or other effective means.
3. Permit the taking of proper precautions to protect food in outdoor bulk fermentation vessels by any effective means, including:
   i. Using protective coverings.
   ii. Controlling areas over and around the vessels to eliminate harборages for pests.
   iii. Checking on a regular basis for pests and pest infestation.
   iv. Skimming the fermentation vessels, as necessary.
(4) Be constructed in such a manner that floors, walls, and ceilings may be adequately cleaned and kept clean and kept in good repair; that drip or condensate from fixtures, ducts and pipes does not contaminate food, food-contact surfaces, food-packaging materials; and that aisles or working spaces are provided between equipment and walls and are adequately unobstructed and of adequate width to permit employees to perform their duties and to protect against contaminating food or food-contact surfaces with clothing or personal contact.

(5) Provide adequate lighting in hand-washing areas, dressing and locker rooms, and toilet rooms and in all areas where food is examined, processed, or stored and where equipment utensils are cleaned; and provide safety-type light bulbs, fixtures, skylights, or other glass suspended over exposed food in any step of preparation or otherwise protect against food contamination in case of glass breakage.

(6) Provide adequate ventilation or control equipment to minimize odors and vapors (including steam and noxious fumes) in areas where they may contaminate food; and locate operate fans and other air-blowing equipment in a manner that minimizes the potential for contaminating food, food-packaging materials, and food-contact surfaces.

(7) Provide, where necessary, adequate screening or other protection against pests.

§ 110.35 Sanitary operations.

(a) General maintenance. Buildings, fixtures, and other physical facilities of the plant shall be maintained in a sanitary condition and shall be kept in repair sufficient to prevent food from becoming adulterated within the meaning of the act. Cleaning and sanitizing of utensils and equipment shall be conducted in a manner that protects against contamination of food, food-contact surfaces, or food-packaging materials.

(b) Substances used in cleaning and sanitizing; storage of toxic materials. (1) Cleaning compounds and sanitizing agents used in cleaning and sanitizing procedures shall be free from undesirable microorganisms and shall be safe and adequate under the conditions of use. Compliance with this requirement may be verified by any effective means including purchase of these substances under a supplier’s guarantee or certification, or examination of these substances for contamination. Only the following toxic materials may be used or stored in a plant where food is processed or exposed:

(i) Those required to maintain clean and sanitary conditions;

(ii) Those necessary for use in laboratory testing procedures;

(iii) Those necessary for plant and equipment maintenance and operation; and

(iv) Those necessary for use in the plant’s operations.
(2) Toxic cleaning compounds, sanitizing agents, and pesticide chemicals shall be identified, held, and stored in a manner that protects against contamination of food, food-contact surfaces, or food-packaging materials. All relevant regulations promulgated by other Federal, State, and local government agencies for the application, use, or holding of these products should be followed.

(c) **Pest control.** No pests shall be allowed in any area of a food plant. Guard or guide dogs may be allowed in some areas of a plant if the presence of the dogs is unlikely to result in contamination of food, food-contact surfaces, or food-packaging materials. Effective measures shall be taken to exclude pests from the processing areas and to protect against the contamination of food on the premises by pests. The use of insecticides or rodenticides is permitted only under precautions and restrictions that will protect against the contamination of food, food-contact surfaces, and food-packaging materials.

(d) **Sanitation of food-contact surfaces.** All food-contact surfaces, including utensils and food-contact surfaces of equipment, shall be cleaned as frequently as necessary to protect against contamination of food.

1. Food-contact surfaces used for manufacturing or holding low-moisture food shall be in a dry, sanitary condition at the time of use. When the surfaces are wet-cleaned, they shall, when necessary, be sanitized and thoroughly dried before subsequent use.

2. In wet processing, when cleaning is necessary to protect against the introduction of microorganisms into food, all food-contact surfaces shall be cleaned and sanitized before use and after any interruption during which the food-contact surfaces may have become contaminated. Where equipment and utensils are used in a continuous production operation the utensils and food-contact surfaces of the equipment shall be cleaned and sanitized as necessary.

3. Non-food-contact surfaces of equipment used in the operation of food plants should be cleaned as frequently as necessary to protect against contamination of food.

4. Single-service articles (such as utensils intended for one-time use, paper cups, and paper towels) should be stored in appropriate containers and shall be handled, dispensed, used, and disposed of in a manner that protects against contamination of food or food-contact surfaces.

5. Sanitizing agents shall be adequate and safe under conditions of use. Any facility, procedure, or machine is acceptable for cleaning and sanitizing equipment and utensils if it is established that the facility, procedure, or machine will routinely render equipment and utensils clean and provide adequate cleaning and sanitizing treatment.

(e) **Storage and handling of cleaned portable equipment and utensils.** Cleaned and sanitized portable equipment with food-contact surfaces and utensils should be stored in a location and manner that protects food-contact surfaces from contamination.

[51 FR 24475, June 19, 1986, as amended at 54 FR 24892, June 12, 1989]
§ 110.37 Sanitary facilities and controls.

Each plant shall be equipped with adequate sanitary facilities and accommodations including, but not limited to:

(a) Water supply. The water supply shall be sufficient for the operations intended and shall be derived from an adequate source. Any water that contacts food or food-contact surfaces shall be safe and of adequate sanitary quality. Running water at a suitable temperature, and under pressure as needed, shall be provided in all areas where required for the processing of food, for the cleaning of equipment, utensils, and food-packaging materials, or for employee sanitary facilities.

(b) Plumbing. Plumbing shall be of adequate size and design and adequately installed and maintained to:
   (1) Carry sufficient quantities of water to required locations throughout the plant.
   (2) Properly convey sewage and liquid disposable waste from the plant.
   (3) Avoid constituting a source of contamination to food, water supplies, equipment, or utensils or creating an unsanitary condition.
   (4) Provide adequate floor drainage in all areas where floors are subject to flooding-type cleaning or where normal operations release or discharge water or other liquid waste on the floor.
   (5) Provide that there is not backflow from, or cross-connection between, piping systems that discharge waste water or sewage and piping systems that carry water for food or food manufacturing.

(c) Sewage disposal. Sewage disposal shall be made into an adequate sewerage system or disposed of through other adequate means.

(d) Toilet facilities. Each plant shall provide its employees with adequate, readily accessible toilet facilities. Compliance with this requirement may be accomplished by:
   (1) Maintaining the facilities in a sanitary condition.
   (2) Keeping the facilities in good repair at all times.
   (3) Providing self-closing doors.
   (4) Providing doors that do not open into areas where food is exposed to airborne contamination, except where alternate means have been taken to protect against such contamination (such as double doors or positive air-flow systems).

(e) Hand-washing facilities. Hand-washing facilities shall be adequate and convenient and be furnished with running water at a suitable temperature. Compliance with this requirement may be accomplished by providing:
   (1) Hand-washing and, where appropriate, hand-sanitizing facilities at each location in the plant where good sanitary practices require employees to wash and/or sanitize their hands.
   (2) Effective hand-cleaning and sanitizing preparations.
   (3) Sanitary towel service or suitable drying devices.
(4) Devices or fixtures, such as water control valves, so designed and constructed to protect against recontamination of clean, sanitized hands.

(5) Readily understandable signs directing employees handling unprotected food, unprotected food-packaging materials, of food-contact surfaces to wash and, where appropriate, sanitize their hands before they start work, after each absence from post of duty, and when their hands may have become soiled or contaminated. These signs may be posted in the processing room(s) and in all other areas where employees may handle such food, materials, or surfaces.

(6) Refuse receptacles that are constructed and maintained in a manner that protects against contamination of food.

(f) Rubbish and offal disposal. Rubbish and any offal shall be so conveyed, stored, and disposed of as to minimize the development of odor, minimize the potential for the waste becoming an attractant and harborage or breeding place for pests, and protect against contamination of food, food-contact surfaces, water supplies, and ground surfaces.

Subpart C – Equipment

§ 110.40 Equipment and utensils.

(a) All plant equipment and utensils shall be so designed and of such material and workmanship as to be adequately cleanable, and shall be properly maintained. The design, construction, and use of equipment and utensils shall preclude the adulteration of food with lubricants, fuel, metal fragments, contaminated water, or any other contaminants. All equipment should be so installed and maintained as to facilitate the cleaning of the equipment and of all adjacent spaces. Food-contact surfaces shall be corrosion-resistant when in contact with food. They shall be made of nontoxic materials and designed to withstand the environment of their intended use and the action of food, and, if applicable, cleaning compounds and sanitizing agents. Food-contact surfaces shall be maintained to protect food from being contaminated by any source, including unlawful indirect food additives.

(b) Seams on food-contact surfaces shall be smoothly bonded or maintained so as to minimize accumulation of food particles, dirt, and organic matter and thus minimize the opportunity for growth of microorganisms.

(c) Equipment that is in the manufacturing or food-handling area and that does not come into contact with food shall be so constructed that it can be kept in a clean condition.

(d) Holding, conveying, and manufacturing systems, including gravimetric, pneumatic, closed, and automated systems, shall be of a design and construction that enables them to be maintained in an appropriate sanitary condition.
(e) Each freezer and cold storage compartment used to store and hold food capable of supporting growth of microorganisms shall be fitted with an indicating thermometer, temperature-measuring device, or temperature-recording device so installed as to show the temperature accurately within the compartment, and should be fitted with an automatic control for regulating temperature or with an automatic alarm system to indicate a significant temperature change in a manual operation.

(f) Instruments and controls used for measuring, regulating, or recording temperatures, pH, acidity, water activity, or other conditions that control or prevent the growth of undesirable microorganisms in food shall be accurate and adequately maintained, and adequate in number for their designated uses.

(g) Compressed air or other gases mechanically introduced into food or used to clean food-contact surfaces or equipment shall be treated in such a way that food is not contaminated with unlawful indirect food additives.

Subpart D – [Reserved]

Subpart E – Production and Process Controls

§ 110.80 Processes and controls.

All operations in the receiving, inspecting, transporting, segregating, preparing, manufacturing, packaging, and storing of food shall be conducted in accordance with adequate sanitation principles. Appropriate quality control operations shall be employed to ensure that food is suitable for human consumption and that food-packaging materials are safe and suitable. Overall sanitation of the plant shall be under the supervision of one or more competent individuals assigned responsibility for this function. All reasonable precautions shall be taken to ensure that production procedures do not contribute contamination from any source. Chemical, microbial, or extraneous-material testing procedures shall be used where necessary to identify sanitation failures or possible food contamination. All food that has become contaminated to the extent that it is adulterated within the meaning of the act shall be rejected, or if permissible, treated or processed to eliminate the contamination.

(a) Raw materials and other ingredients.

(1) Raw materials and other ingredients shall be inspected and segregated or otherwise handled as necessary to ascertain that they are clean and suitable for processing into food and shall be stored under conditions that will protect against contamination and minimize deterioration. Raw materials shall be washed or cleaned as necessary to remove soil or other contamination. Water used for washing, rinsing, or conveying food shall be safe and of adequate sanitary quality. Water may be reused for washing, rinsing, or conveying food if it does not increase the level of contamination of the food. Containers and carriers of raw materials should be inspected on receipt to ensure that their condition has not contributed to the contamination or deterioration of food.
(2) Raw materials and other ingredients shall either not contain levels of microorganisms that may produce food poisoning or other disease in humans, or they shall be pasteurized or otherwise treated during manufacturing operations so that they no longer contain levels that would cause the product to be adulterated within the meaning of the act. Compliance with this requirement may be verified by any effective means, including purchasing raw materials and other ingredients under a supplier’s guarantee or certification.

(3) Raw materials and other ingredients susceptible to contamination with aflatoxin or other natural toxins shall comply with current Food and Drug Administration regulations and action levels for poisonous or deleterious substances before these materials or ingredients are incorporated into finished food. Compliance with this requirement may be accomplished by purchasing raw materials and other ingredients under a supplier's guarantee or certification, or may be verified by analyzing these materials and ingredients for aflatoxins and other natural toxins.

(4) Raw materials, other ingredients, and rework susceptible to contamination with pests, undesirable microorganisms, or extraneous material shall comply with applicable Food and Drug Administration regulations and defect action levels for natural or unavoidable defects if a manufacturer wishes to use the materials in manufacturing food. Compliance with this requirement may be verified by any effective means, including purchasing the materials under a supplier's guarantee or certification, or examination of these materials for contamination.

(5) Raw materials, other ingredients, and rework shall be held in bulk, or in containers designed and constructed so as to protect against contamination and shall be held at such temperature and relative humidity and in such a manner as to prevent the food from becoming adulterated within the meaning of the act. Material scheduled for rework shall be identified as such.

(6) Frozen raw materials and other ingredients shall be kept frozen. If thawing is required prior to use, it shall be done in a manner that prevents the raw materials and other ingredients from becoming adulterated within the meaning of the act.

(7) Liquid or dry raw materials and other ingredients received and stored in bulk form shall be held in a manner that protects against contamination.

(b) **Manufacturing operations.**

(1) Equipment and utensils and finished food containers shall be maintained in an acceptable condition through appropriate cleaning and sanitizing, as necessary. Insofar as necessary, equipment shall be taken apart for thorough cleaning.

(2) All food manufacturing, including packaging and storage, shall be conducted under such conditions and controls as are necessary to minimize the potential for the growth of microorganisms, or for the
Appendix 3

contamination of food. One way to comply with this requirement is careful monitoring of physical factors such as time, temperature, humidity, $a_w$, pH, pressure, flow rate, and manufacturing operations such as freezing, dehydration, heat processing, acidification, and refrigeration to ensure that mechanical breakdowns, time delays, temperature fluctuations, and other factors do not contribute to the decomposition or contamination of food.

(3) Food that can support the rapid growth of undesirable microorganisms, particularly those of public health significance, shall be held in a manner that prevents the food from becoming adulterated within the meaning of the act. Compliance with this requirement may be accomplished by any effective means, including:

(i) Maintaining refrigerated foods at $45^\circ F$ ($7.2^\circ C$) or below as appropriate for the particular food involved.
(ii) Maintaining frozen foods in a frozen state.
(iii) Maintaining hot foods at $140^\circ F$ ($60^\circ C$) or above.
(iv) Heat treating acid or acidified foods to destroy mesophilic microorganisms when those foods are to be held in hermetically sealed containers at ambient temperatures.

(4) Measures such as sterilizing, irradiating, pasteurizing, freezing, refrigerating, controlling pH or controlling $a_w$ that are taken to destroy or prevent the growth of undesirable microorganisms, particularly those of public health significance, shall be adequate under the conditions of manufacture, handling, and distribution to prevent food from being adulterated within the meaning of the act.

(5) Work-in-process shall be handled in a manner that protects against contamination.

(6) Effective measures shall be taken to protect finished food from contamination by raw materials, other ingredients, or refuse. When raw materials, other ingredients, or refuse are unprotected, they shall not be handled simultaneously in a receiving, loading, or shipping area if that handling could result in contaminated food. Food transported by conveyor shall be protected against contamination as necessary.

(7) Equipment, containers, and utensils used to convey, hold, or store raw materials, work-in-process, rework, or food shall be constructed, handled, and maintained during manufacturing or storage in a manner that protects against contamination.

(8) Effective measures shall be taken to protect against the inclusion of metal or other extraneous material in food. Compliance with this requirement may be accomplished by using sieves, traps, magnets, electronic metal detectors, or other suitable effective means.

(9) Food, raw materials, and other ingredients that are adulterated within the meaning of the act shall be disposed of in a manner that protects against the contamination of other food. If the adulterated food is capable of being reconditioned, it shall be reconditioned using a method that has been proven to be effective or it shall be reexamined and found not to be adulterated within the meaning of the act before being incorporated into other food.
(10) Mechanical manufacturing steps such as washing, peeling, trimming, cutting, sorting and inspecting, mashing, dewatering, cooling, shredding, extruding, drying, whipping, defatting, and forming shall be performed so as to protect food against contamination. Compliance with this requirement may be accomplished by providing adequate physical protection of food from contaminants that may drip, drain, or be drawn into the food. Protection may be provided by adequate cleaning and sanitizing of all food-contact surfaces, and by using time and temperature controls at and between each manufacturing step.

(11) Heat blanching, when required in the preparation of food, should be effected by heating the food to the required temperature, holding it at this temperature for the required time, and then either rapidly cooling the food or passing it to subsequent manufacturing without delay. Thermophilic growth and contamination in blanchers should be minimized by the use of adequate operating temperatures and by periodic cleaning. Where the blanched food is washed prior to filling, water used shall be safe and of adequate sanitary quality.

(12) Batters, breading, sauces, gravies, dressings, and other similar preparations shall be treated or maintained in such a manner that they are protected against contamination. Compliance with this requirement may be accomplished by any effective means, including one or more of the following:
   (i) Using ingredients free of contamination.
   (ii) Employing adequate heat processes where applicable.
   (iii) Using adequate time and temperature controls.
   (iv) Providing adequate physical protection of components from contaminants that may drip, drain, or be drawn into them.
   (v) Cooling to an adequate temperature during manufacturing.
   (vi) Disposing of batters at appropriate intervals to protect against the growth of microorganisms.

(13) Filling, assembling, packaging, and other operations shall be performed in such a way that the food is protected against contamination. Compliance with this requirement may be accomplished by any effective means, including:
   (i) Use of a quality control operation in which the critical control points are identified and controlled during manufacturing.
   (ii) Adequate cleaning and sanitizing of all food-contact surfaces and food containers.
   (iii) Using materials for food containers and food-packaging materials that are safe and suitable, as defined in Sec. 130.3(d) of this chapter.
   (iv) Providing physical protection from contamination, particularly airborne contamination.
   (v) Using sanitary handling procedures.
(14) Food such as, but not limited to, dry mixes, nuts, intermediate moisture food, and dehydrated food, that relies on the control of $a_w$ for preventing the growth of undesirable microorganisms shall be processed to and maintained at a safe moisture level. Compliance with this requirement may be accomplished by any effective means, including employment of one or more of the following practices:

(i) Monitoring the $a_w$ of food.

(ii) Controlling the soluble solids-water ratio in finished food.

(iii) Protecting finished food from moisture pickup, by use of a moisture barrier or by other means, so that the $a_w$ of the food does not increase to an unsafe level.

(15) Food such as, but not limited to, acid and acidified food, that relies principally on the control of pH for preventing the growth of undesirable microorganisms shall be monitored and maintained at a pH of 4.6 or below. Compliance with this requirement may be accomplished by any effective means, including employment of one or more of the following practices:

(i) Monitoring the pH of raw materials, food in process, and finished food.

(ii) Controlling the amount of acid or acidified food added to low-acid food.

(16) When ice is used in contact with food, it shall be made from water that is safe and of adequate sanitary quality, and shall be used only if it has been manufactured in accordance with current good manufacturing practice as outlined in this part.

(17) Food-manufacturing areas and equipment used for manufacturing human food should not be used to manufacture nonhuman food-grade animal feed or inedible products, unless there is no reasonable possibility for the contamination of the human food.


§ 110.93 Warehousing and distribution.

Storage and transportation of finished food shall be under conditions that will protect food against physical, chemical, and microbial contamination as well as against deterioration of the food and the container.

Subpart F – [Reserved]

Subpart G – Defect Action Levels

§ 110.110 Natural or unavoidable defects in food for human use that present no health hazard.

(a) Some foods, even when produced under current good manufacturing practice, contain natural or unavoidable defects that at low levels are not hazardous to health. The Food and Drug Administration establishes maximum levels for these defects in foods produced under current good manufacturing practice and uses these levels in deciding whether to recommend regulatory action.
(b) Defect action levels are established for foods whenever it is necessary and feasible to do so. These levels are subject to change upon the development of new technology or the availability of new information.

(c) Compliance with defect action levels does not excuse violation of the requirement in section 402(a)(4) of the act that food not be prepared, packed, or held under unsanitary conditions or the requirements in this part that food manufacturers, distributors, and holders shall observe current good manufacturing practice. Evidence indicating that such a violation exists causes the food to be adulterated within the meaning of the act, even though the amounts of natural or unavoidable defects are lower than the currently established defect action levels. The manufacturer, distributor, and holder of food shall at all times utilize quality control operations that reduce natural or unavoidable defects to the lowest level currently feasible.

(d) The mixing of a food containing defects above the current defect action level with another lot of food is not permitted and renders the final food adulterated within the meaning of the act, regardless of the defect level of the final food.

(e) A compilation of the current defect action levels for natural or unavoidable defects in food for human use that present no health hazard may be obtained upon request from the Center for Food Safety and Applied Nutrition (HFS-565), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740.

Traditional Approach for Conducting Hazard Analysis

Fresh mahi-mahi Example

Product Description: Fresh wild mahi-mahi fillets

Fishery Product Market Name: Mahi-mahi (Coryphaena species)

Source of Fishery Product: From other processors, received in ice

Methods of Packaging, Distribution and Storage: Air packed, stored and distributed on ice

Intended Use and Consumer: To be cooked and consumed by the general public

Description of Process

Receive fresh fish – Fresh wild caught mahi-mahi (Coryphaena species, not aquacultured) fillets are received from several domestic suppliers. Delivery truck transit times range from 2 to 8 hours. Tubs or other containers of mahi-mahi fillets are received along with other fresh seafood products packed in ice and delivered by refrigerated truck. After receipt, products are re-iced if necessary and moved into refrigerated storage.

Refrigerated storage – Individual mahi-mahi fillets are completely buried in ice and stored in a refrigerated cooler until needed.

Trim – Individual tubs or containers of mahi-mahi fillets are removed from the cooler as needed to pack customer orders. Fillets are trimmed by hand with knives if necessary to meet customer specifications. Trimming is completed in 30 minutes or less.
**Weigh/Pack/Label** – Per customer order, mahi-mahi fillets are weighed, packed into containers, and each container is labeled with a handwritten or printed label that contains the market name of the species of fish or shellfish that it contains. Individual containers are completely surrounded by ice and assembled into master cartons for each customer order. The weigh/pack/label step is completed in 30 minutes or less.

**Finished product storage** – Iced mahi-mahi containers in master cartons that contain each customer’s order are placed back into refrigerated storage until it is moved directly to refrigerated trucks for delivery to retail or restaurant customers.

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![Fresh Mahi-mahi Fillets Process Flow Chart](image-url)
<table>
<thead>
<tr>
<th>Processing Step</th>
<th>(1) Biological, chemical, and physical food safety hazards</th>
<th>(2)</th>
<th>(3) Is the potential food safety hazard significant? (Yes or No)</th>
<th>(4) Justify the decision that you made in column 3</th>
<th>(5) What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard?</th>
<th>(6) Is this step a Critical Control Point? (Yes or No)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receiving fresh fillets</td>
<td>BIOLOGICAL Pathogen contamination</td>
<td>No</td>
<td>Product will be fully cooked prior to consumption</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pathogen growth</td>
<td>No</td>
<td>Product will be fully cooked prior to consumption</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CHEMICAL Histamine</td>
<td>Yes</td>
<td>Time/temperature abuse during transit could cause histamine to form in fillets</td>
<td>Mahi-mahi fillets are shipped in containers buried in ice</td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Food Allergens</td>
<td>Yes</td>
<td>Fish is a food allergen</td>
<td>Fillets will be labeled with market name at weigh/pack/label step</td>
<td>No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Refrigerated Storage</td>
<td>BIOLOGICAL Bacterial pathogen growth</td>
<td>No</td>
<td>Product will be fully cooked prior to consumption</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CHEMICAL Histamine</td>
<td>Yes</td>
<td>Time/temperature abused during storage could cause histamine formation</td>
<td>Mahi-mahi fillets are buried in ice &amp; stored in a refrigerated cooler</td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Food Allergens</td>
<td>No</td>
<td>Fillets will be labeled with market name at weigh/pack/label step</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PHYSICAL None</td>
<td>No</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processing Step</td>
<td>[2] List all potential biological, chemical, and physical food safety hazards that could be associated with this product and process.</td>
<td>[3] Is the potential food safety hazard significant (introduced, enhanced or eliminated) at this step? (Yes or No)</td>
<td>[4] Justify the decision that you made in column 3</td>
<td>[5] What control measure(s) can be applied to prevent, eliminate or reduce this significant hazard?</td>
<td>[6] Is this step a Critical Control Point? (Yes or No)</td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------------------------</td>
<td>------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------</td>
<td>-------------------------------------------------</td>
<td></td>
</tr>
</tbody>
</table>
| Trimming            | BIOLOGICAL Bacterial Pathogen Contamination  
                      | No  
                      | Controlled by SSOP  |                                                                 |  
                      |  
                      |                                                                                       |
|                     | CHEMICAL Histamine  
                      | No  
                      | Not likely to occur. Time at this and weigh/pack/label step is 30 minutes or less  |                                                                 |  
                      |  
                      |                                                                                       |
|                     | Food Allergens  
                      | No  
                      | Fillets will be labeled with market name at weigh/pack/label step  |                                                                 |  
                      |  
                      |                                                                                       |
|                     | PHYSICAL Metal Contamination  
                      | No  
                      | Fillet knives are not likely to chip and contaminate product with metal  |                                                                 |  
                      |  
                      |                                                                                       |
| Weight/Pack/Label   | BIOLOGICAL Bacterial Pathogen Contamination  
                      | No  
                      | Controlled by SSOP  |                                                                 |  
                      |  
                      |                                                                                       |
|                     | CHEMICAL Histamine  
                      | No  
                      | Not likely to occur. Time at this and weigh/pack/label step is 30 minutes or less  |                                                                 |  
                      |  
                      |                                                                                       |
|                     | Food Allergens  
                      | Yes  
                      | Fish is a food allergen  
                      | Fillets will be labeled with market name at this step  | Yes  
                      |  
                      |                                                                                       |
|                     | PHYSICAL None  
                      |  
                      |                                                                 |  
                      |  
                      |  
                      |                                                                                       |
| Finished Product    | BIOLOGICAL Bacterial Pathogen Growth  
                      | No  
                      | Product will be fully cooked prior to consumption  |                                                                 |  
                      |  
                      |                                                                                       |
| Refrigerated Storage| CHEMICAL Histamine  
                      | Yes  
                      | Time/temperature abused during storage could cause histamine formation  
                      | Mahi-mahi fillets will be buried in ice & stored in a refrigerated cooler  | Yes  
                      |  
                      |                                                                                       |
|                     | PHYSICAL None  
<pre><code>                  |  
                  |                                                                 |  
                  |  
                  |  
                  |                                                                                       |
</code></pre>
<table>
<thead>
<tr>
<th>Critical Control Point (CCP)</th>
<th>Significant Hazard(s)</th>
<th>Critical Limits for each Control Measure</th>
<th>Monitoring</th>
<th>Corrective Action</th>
<th>Verification</th>
<th>Records</th>
</tr>
</thead>
<tbody>
<tr>
<td>Receving</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice at receipt.</td>
<td>Adequacy of ice surrounding containers of fillets at delivery.</td>
<td>Visual check of adequacy of ice in a representative number of containers in each delivery.</td>
<td>Every Delivery. Receiving Manager.</td>
<td>Weekly review of Receiving Log (Monitoring record) and Corrective Action and Verification records.</td>
</tr>
<tr>
<td>Refrigerated Storage</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td>Adequacy of ice surrounding containers of fillets</td>
<td>Visual check of adequacy of ice in a representative number of containers in cooler storage.</td>
<td>At the beginning and end of the work day. Cooler Manager.</td>
<td>Weekly review of Cooler Ice Log (Monitoring Record) and Corrective action and Verification records.</td>
</tr>
<tr>
<td>Critical Control Point (CCP)</td>
<td>Significant Hazard(s)</td>
<td>Critical Limits for each Control Measure</td>
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</tr>
<tr>
<td>Weigh/Pack/Label</td>
<td>Food Allergens</td>
<td>All finished product containers will be labeled with the correct market name of the fish.</td>
<td>The market name on each container of finished product</td>
<td>Visual check of a representative number of containers and their label</td>
<td>If a container is improperly labeled, then: segregate it and properly label it before the customer order is placed in the finished product cooler and modify labeling procedure and conduct training as necessary to ensure that all products are properly identified.</td>
<td>Weekly review of Packing Room Log (Monitoring record) and Corrective action and Verification records</td>
</tr>
<tr>
<td>Finished Product Refrigerated Storage</td>
<td>Histamine</td>
<td>Mahi-mahi fillets are completely surrounded with ice throughout storage time.</td>
<td>Adequacy of ice surrounding containers of fillets</td>
<td>Visual check of representative number of containers in cooler storage</td>
<td>If finished product containers do not have adequate ice, then: chill and hold the product until it can be evaluated based on its total time and temperature exposure including exposures during prior processing operations, and determine if there is a problem with the cooler and fix it.</td>
<td>Weekly review of Cooler Ice Log (Monitoring record), Corrective action and Verification records</td>
</tr>
</tbody>
</table>

Firm Name: XYZ Seafood Company
Product: Fresh mahi-mahi fillets
Firm Address: 238 Coastal Lane, Happy Beach, XX
Method of Storage and Distribution: Stored and distributed buried in ice
Intended Use and Consumer: To be cooked and consumed by the general public
Signature: ____________________________
Print name: ____________________________
Date: ____________________________
Also Available from the National Seafood HACCP Alliance

**Guidance for Industry: Fish and Fishery Products Hazards and Controls Guidance (Fourth Edition)**

The newly revised FDA Hazards Guide will assist seafood industry compliance with FDA regulations that cover domestic and imported seafood. Key updates include post-harvest treatment information for pathogenic bacteria in shellfish; time and temperature adjustments to control histamine formation and pathogenic bacteria food safety hazards; consistency with changes in statutes regulations, tolerance and action levels for food additives, aquaculture drug approvals, natural toxins, chemicals and pesticides; species hazard identification; and listing potential public health consequences of seafood safety hazards. This is a companion document to the Seafood HACCP Training Curriculum Manual (SGR 127). $25.

**HACCP: Programa de Capacitación en Análisis de Peligros y Puntos Críticos de Control**

Este manual de enseñanza para educación HACCP incluye la identificación de peligros, la regulación HACCP del U.S. FDA y desarrollo de plan HACCP. Este es un manual complementario a la “Guía de Peligros y Controles para Pescados y Productos Pesqueros” del U.S. FDA, y debe utilizarse en conjunto con la misma para desarrollar y evaluar planes HACCP. $25.

**Sanitation Control Procedures for Processing Fish and Fishery Products**

This course is intended to assist the seafood industry in developing and implementing sanitation control procedures as mandated by the Food and Drug Administration (FDA). These mandates require seafood processors to monitor sanitary control procedures used during processing in order to show their compliance with approved sanitary conditions and practices. Likewise, seafood importers must verify that the seafood imported was processed in accordance with the same FDA-mandated HACCP requirements that include sanitation procedure monitoring and records. $25.

**Curso sobre Prócedimientos de Control Sanitario para el Procesamiento de Pescados y Mariscos**

Esta guía es el manual de entrenamiento para el curso dictado con el propósito de asistir a la industria pesquera en el desarrollo e implementación de Procedimientos de Control Sanitarios como requeridos por la Administración de Drogas y Alimentos de Estados Unidos de América (US FDA). Estos mandatos requieren que los procesadores de productos pesqueros monitoreen los procedimientos de control sanitarios utilizados durante el proceso para documentar el cumplimiento con las prácticas y condiciones sanitarias aprobadas. Igualmente, los importadores de productos pesqueros deben verificar que los productos importados fueron procesados bajo los mismos requisitos HACCP de la US FDA que incluyen el monitoreo y registros de los procesos sanitarios. $25.

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