WORLD ORGANISATION FOR ANIMAL HEALTH

Protecting animals, preserving our future

AQUATIC ANIMAL HEALTH CODE

Twentieth Edition, 2017
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**Index**
The OIE Aquatic Animal Health Code (the Aquatic Code) sets out standards for the improvement of aquatic animal health and welfare of farmed fish worldwide, and for safe international trade in aquatic animals (amphibians, crustaceans, fish and molluscs) and their products. The health measures in the Aquatic Code should be used by the Competent Authorities of importing and exporting countries for early detection, reporting and control of agents pathogenic to aquatic animals and to prevent their transfer via international trade in aquatic animals and their products, while avoiding unjustified sanitary barriers to trade.

The standards in the Aquatic Code have been formally adopted by the World Assembly of OIE Delegates, which constitutes the organisation's highest decision-making body. This 20th edition incorporates modifications to the Aquatic Code agreed at the 85th General Session in May 2017.

This edition includes the following updates:

- revisions to several definitions in the glossary;
- Chapter 1.2. 'Criteria for listing aquatic animal diseases' has been extensively amended to align with the corresponding chapter in the OIE Terrestrial Animal Health Code;
- a new disease, Batrachochytrium salamandrivorans, has been added to Chapter 1.3. 'Diseases listed by the OIE' and some disease names have been amended;
- minor amendments have been made in Chapters 4.3. 'Disinfection of aquaculture establishments and equipment', 4.4. 'Recommendations for surface disinfection of salmonid eggs' and 5.1. 'General obligations related to certification';
- a number of horizontal amendments were made in all crustacean disease-specific chapters, to improve readability. In addition, the list of susceptible species in Article X.X.2. in Chapters 9.2., 9.3., 9.4., 9.5., 9.6. and 9.8. was reviewed and amended, where relevant, after consideration of the application of the 'Criteria for listing species as susceptible to infection with a specific pathogen' (Chapter 1.5.);
- a new chapter on acute hepatopancreatic necrosis disease (Chapter 9.1.) has been added;
- Article X.X.8. in all disease-specific chapters was revised to more adequately describe the requirements for the importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from disease X;
- the year that a chapter was first adopted and the year of last revision are noted at the end of each chapter. In this regard the OIE has made every endeavour to ensure the accuracy of this information based on our historical records.

The development of these standards and recommendations is the result of the ongoing work by the OIE Aquatic Animal Health Standards Commission (the Aquatic Animals Commission). This Commission, which comprises six elected members, meets twice yearly to address its work programme. This Commission draws upon the expertise of internationally renowned specialists to prepare draft texts for new articles of the Aquatic Code and to revise existing articles. The views of OIE National Delegates are routinely sought through the twice yearly circulation of new or revised texts. The Aquatic Animals Commission collaborates closely with other Specialist Commissions of the OIE, including the Terrestrial Animal Health Standards Commission, the Biological Standards Commission and the Scientific Commission for Animal Diseases, to ensure that the recommendations contained in the Aquatic Code are based upon the latest scientific information.

The World Trade Organization (WTO) Agreement on the Application of Sanitary and Phytosanitary Measures (SPS Agreement) formally recognises the role of the OIE as the international standard setting organisation for animal health and zoonotic diseases. According to the SPS Agreement, WTO Members should align their import requirements with the recommendations in the relevant standards of the OIE. Where there are no OIE recommendations or if the country chooses a level of protection requiring measures more stringent than the standards of the OIE, these should be based on an import risk analysis conducted in accordance with Chapter 2.1. The Aquatic Code is thus a key part of the WTO legal framework for international trade.

The Aquatic Code is published annually in the three official OIE languages (English, French and Spanish). The Aquatic Code may be viewed and downloaded from the OIE Web site at http://www.oie.int.
Foreword

The User's Guide, which follows the foreword, is designed to help Competent Authorities and other interested parties to use the Aquatic Code.

We wish to thank the members of the Aquatic Animals Commission, Delegates and the experts participating in ad hoc Groups and other Specialist Commissions for their expert advice. My thanks go to the staff of the OIE for their dedication in producing this 20th edition of the Aquatic Code.

Dr Monique Eloit  
Director General  
World Organisation for Animal Health

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A. Introduction

1) The OIE Aquatic Animal Health Code (hereafter referred to as the Aquatic Code) provides standards for the improvement of aquatic animal health worldwide. The Aquatic Code also includes standards for the welfare of farmed fish and use of antimicrobial agents in aquatic animals. The purpose of this guide is to advise the Competent Authorities in OIE Member Countries on how to use the Aquatic Code.

2) Competent Authorities should use the standards in the Aquatic Code to develop measures for early detection, internal reporting, notification and control of pathogenic agents in aquatic animals (amphibians, crustaceans, fish and molluscs) and preventing their spread via international trade in aquatic animals and aquatic animal products, while avoiding unjustified sanitary barriers to trade.

3) OIE standards are based on the most recent scientific and technical information. Correctly applied, they protect aquatic animal health during the production and trade in aquatic animals and aquatic animal products as well as the welfare of farmed fish.

4) The absence of chapters, articles or recommendations on particular pathogenic agents or commodities does not preclude the application of appropriate sanitary measures by the Competent Authorities, provided they are based on risk analyses conducted in accordance with the Aquatic Code.

5) The complete text of the Aquatic Code is available on the OIE website and individual chapters may be downloaded from: http://www.oie.int.

B. Aquatic Code content

1) Key terms and expressions used in more than one chapter in the Aquatic Code are defined in the Glossary. The reader should be aware of definitions given in the Glossary when reading and using the Aquatic Code. Defined terms appear in italics. In the online version of the Aquatic Code, a hyperlink leads to the relevant definition.

2) The term ‘(under study)’ is found in some rare instances, with reference to an article or part of an article. This means that this part of the text has not been adopted by the World Assembly of OIE Delegates and the particular provisions are thus not part of the Aquatic Code.

3) The standards in the chapters of Section 1 are designed for the implementation of measures for the surveillance and notification of pathogenic agents. The section includes the criteria for listing aquatic animal diseases, the diseases which are listed by the OIE, procedures for notification to the OIE, and criteria for listing species as susceptible to infection with a specific pathogen.

4) The standards in the chapters of Section 2 are designed to guide the importing country in conducting import risk analysis in the absence of OIE standards. The importing country may also use these standards to justify import measures which are more stringent than existing OIE standards.

5) The standards in the chapters of Section 3 are designed for the establishment, maintenance and evaluation of Aquatic Animal Health Services, including communication. These standards are intended to assist the Competent Authorities of Member Countries to meet their objectives of improving aquatic animal health and welfare of farmed fish, as well as to establish and maintain confidence in their international aquatic animal health certificates.

6) The standards in the chapters of Section 4 are designed for the implementation of measures for the prevention and control of pathogenic agents. Measures in this section include zoning, compartmentalisation, disinfection, contingency planning, fallowing, disposal of aquatic animal waste and control of pathogenic agents in aquatic animal feed.

7) The standards in the chapters of Section 5 are designed for the implementation of general sanitary measures for trade. They address certification and the measures applicable by the exporting, transit and importing countries. A range of model international aquatic animal health certificates is provided to facilitate consistent documentation for international trade.

8) The standards in the chapters of Section 6 are designed to ensure the responsible and prudent use of antimicrobial agents in aquatic animals.

9) The standards in the chapters of Section 7 are designed for the implementation of welfare measures for farmed fish. The standards cover the general principles for welfare of farmed fish, including during transport, stunning and killing for human consumption, and when killing for disease control purposes.
10) The standards in each of the chapters of Sections 8 to 11 are designed to prevent the pathogenic agents of OIE listed diseases from being introduced into an importing country. Each disease chapter includes a list of currently known susceptible species. The standards take into account the nature of the traded commodity, the aquatic animal health status of the exporting country, zone or compartment, and the risk reduction measures applicable to each commodity.

These standards assume that the agent is either not present in the importing country or is the subject of a control or eradication programme. Sections 8 to 11 each relate to amphibian, crustacean, fish and molluscan hosts, respectively.

C. Specific issues

1) Notification

Chapter 1.1. describes Member Countries' obligations under OIE Organic Statutes. Listed diseases, as prescribed in Chapter 1.1., are compulsorily notifiable. Member Countries are encouraged to also provide information to the OIE on other aquatic animal health events of epidemiological significance, including occurrence of emerging diseases.

Chapter 1.2. describes the criteria for the inclusion of a disease listed by the OIE.

Chapter 1.3. specifies the diseases that are listed by the OIE. Diseases are divided into four sections corresponding to amphibian, crustacean, fish and molluscan hosts, respectively.

2) Pathogen differentiation

Some pathogens have one or more variants. Existence of highly pathogenic variants and the need to differentiate them from more benign variants are recognised in the Aquatic Code. When pathogenic agents have strains that are stable, possess characteristics that can be used for diagnostic purposes, and display different levels of pathogenicity, different standards providing protection proportionate to the risk posed by the different strains should be applied. Infection with infectious salmon anaemia virus is the first listed disease for which risk management options based on strain differentiation are provided.

3) Determining the susceptibility of species

The Aquatic Code proposes the use of criteria to assess the susceptibility of host species to the pathogenic agents of diseases listed in the Aquatic Code. This is important in the aquaculture context, given the large number of existing and new aquaculture species.

4) Trade requirements

Aquatic animal health measures related to international trade should be based on OIE standards. A Member Country may authorise the importation of aquatic animals or aquatic animal products into its territory under conditions different from those recommended by the Aquatic Code. To scientifically justify more stringent measures, the importing country should conduct a risk analysis in accordance with OIE standards, as described in Chapter 2.1. Members of the WTO should refer to the Agreement on the Application of Sanitary and Phytosanitary Measures (SPS Agreement).

Chapters 5.1. to 5.3. describe the obligations and ethical responsibilities of importing and exporting countries in international trade. Competent Authorities and all veterinarians and certifying officials directly involved in international trade should be familiar with these chapters. Chapter 5.3. also describes the OIE informal procedure for dispute mediation.

Disease-specific chapters in the Aquatic Code include articles listing the commodities that are considered safe for trade without the imposition of disease-specific sanitary measures, regardless of the status of the exporting country or zone for the pathogenic agent in question. Where such a list is present, importing countries should not require any conditions related to the agent in question with respect to the listed commodities.

5) Trade in aquatic animal commodities

Chapter 5.4. describes the criteria used to assess the safety of aquatic animal commodities.

Based on assessments using criteria in Article 5.4.1., in all disease-specific chapters, point 1 of Article X.X.3. lists aquatic animal commodities that may be imported for any purpose from a country, zone or compartment not declared free from the disease in question. The criteria for inclusion of aquatic animal commodities in point 1 of Article X.X.3. are based on the absence of the pathogenic agent or inactivation of the pathogenic agent by treatment or processing.

Based on assessments using criteria in Article 5.4.2, in all disease-specific chapters, point 1 of Article X.X.12. (for Chapter 10.4. the relevant Article is 10.4.15.) lists aquatic animal commodities for retail trade for human consumption from a country, zone or compartment not declared free from the disease in question. The criteria for inclusion of aquatic animal commodities in point 1 of Article X.X.12. include consideration of the form and presentation of the product, the expected volume of waste tissues generated by the consumer and the likely presence of viable pathogenic agent in the waste.
6) International health certificates

An international aquatic animal health certificate is an official document that the Competent Authority of the exporting country issues in accordance with Chapter 5.1. and Chapter 5.2. It lists aquatic animal health requirements for the exported commodity. The quality of the exporting country's Aquatic Animal Health Services is essential in providing assurances to trading partners regarding the safety of exported aquatic animal commodities. This includes the Aquatic Animal Health Services’ ethical approach to the provision of international health certificates and their history in meeting their notification obligations.

International health certificates underpin international trade and provide assurances to the importing country regarding the health status of the aquatic animal commodities imported. The measures prescribed should take into account the health status of both exporting and importing countries and be based upon the standards in the Aquatic Code.

The following steps should be taken when drafting international aquatic animal health certificates:

a) identify the diseases, from which the importing country is justified in seeking protection because of its own aquatic animal health status. Importing countries should not impose measures in regards to diseases that occur in their own territory but are not subject to official control programmes;

b) for commodities capable of transmitting these diseases through international trade, the importing country should apply the relevant articles in the disease-specific chapters. The application of the articles should be adapted to the disease status of the exporting country, zone or compartment. Such a status should be established in accordance with Article 1.4.6. except when articles of the relevant disease chapter specify otherwise;

c) when preparing international aquatic animal health certificates, the importing country should endeavour to use terms and expressions in accordance with the definitions given in the Glossary. As stated in Article 5.2.3., international aquatic animal health certificates should be kept as simple as possible and should be clearly worded, to avoid misunderstanding of the importing country's requirements;

d) Chapter 5.10. provides, as further guidance to Member Countries, model health certificates that should be used as a baseline.

7) Guidance notes for importers and exporters

It is recommended that Competent Authorities prepare 'guidance notes' to assist importers and exporters to understand trade requirements. These notes should identify and explain the trade conditions, including the measures to be applied before and after export and during transport and unloading, and the relevant legal obligations and operational procedures. The guidance notes should advise on all details to be included in the health certification accompanying the consignment to its destination. Exporters should also be reminded of the International Air Transport Association rules governing air transport of aquatic animals and aquatic animal products.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2015.
GLOSSARY

For the purpose of the Aquatic Code:

**ANTIMICROBIAL AGENT**

means a naturally occurring, semi-synthetic or synthetic substance that at in vivo concentrations exhibits antimicrobial activity (kill or inhibit the growth of microorganisms). Anthelmintics and substances classed as disinfectants or antiseptics are excluded from this definition.

**AQUACULTURE**

means the farming of aquatic animals with some sort of intervention in the rearing process to enhance production, such as regular stocking, feeding, protection from predators, etc.

**AQUACULTURE ESTABLISHMENT**

means an establishment in which amphibians, fish, molluscs or crustaceans for breeding, stocking or sale are raised or kept.

**AQUATIC ANIMAL HEALTH PROFESSIONAL**

means a person who, for the purposes of the Aquatic Code, is authorised by the Competent Authority to carry out certain designated tasks in a territory and has the appropriate qualifications and training to perform the designated tasks.

**AQUATIC ANIMAL HEALTH SERVICES**

means the governmental and non-governmental organisations that implement animal health and welfare measures and other standards and recommendations in the Aquatic Code in the territory. The Aquatic Animal Health Services are under the overall control and direction of the Competent Authority. Private sector organisations, veterinarians or aquatic animal health professionals are normally accredited or approved by the Competent Authority to deliver the delegated functions.

**AQUATIC ANIMAL HEALTH STATUS**

means the status of a country, zone or compartment with respect to an aquatic animal disease in accordance with the criteria listed in the relevant chapter of the Aquatic Code dealing with the disease.

**AQUATIC ANIMAL PRODUCTS**

means non-viable aquatic animals and products from aquatic animals.

**AQUATIC ANIMALS**

means all viable life stages (including eggs and gametes) of fish, molluscs, crustaceans and amphibians originating from aquaculture establishments or from the wild.

**Aquatic Code**

means the OIE Aquatic Animal Health Code.

**Aquatic Manual**

means the OIE Manual of Diagnostic Tests for Aquatic Animals.

**BASIC BIOSECURITY CONDITIONS**

means a set of conditions applying to a particular disease, and a particular zone or country, required to ensure adequate disease security, such as:

a) the disease, including suspicion of the disease, is compulsorily notifiable to the Competent Authority; and

b) an early detection system is in place within the zone or country; and

c) import requirements to prevent the introduction of disease into the country or zone, as outlined in the Aquatic Code, are in place.
Glossary

BIAS

means a tendency of an estimate to differ in a non-random fashion from the true value of a population parameter.

BIOLOGICAL PRODUCTS

means:

a) biological reagents for use in the diagnosis of certain diseases;
b) sera for use in the prevention and treatment of certain diseases;
c) inactivated or modified vaccines for use in preventive vaccination against certain diseases;
d) genetic material of pathogenic agents;
e) endocrine tissues from fish or used in fish.

BIOSECURITY

means a set of management and physical measures designed to reduce the risk of introduction, establishment and spread of pathogenic agents to, from and within an aquatic animal population.

BIOSECURITY PLAN

means a plan that identifies significant potential pathways for the introduction and spread of disease in a zone or compartment, and describes the measures which are being, or will be, applied to mitigate the risks to introduce and spread disease, taking into consideration the recommendations in the Aquatic Code. The plan should also describe how these measures are audited, with respect to both their implementation and their targeting, to ensure that the risks are regularly re-assessed and the measures adjusted accordingly.

CASE

means an individual aquatic animal infected by a pathogenic agent, with or without clinical signs.

CASE DEFINITION

is a set of criteria used to distinguish a case animal or an epidemiological unit from a non-case.

CERTIFYING OFFICIAL

means a person authorised by the Competent Authority to sign health certificates for aquatic animals.

COMMODITY

means aquatic animals, aquatic animal products, biological products and pathological material.

COMPARTMENT

means one or more aquaculture establishments under a common biosecurity management system containing an aquatic animal population with a distinct health status with respect to a specific disease or diseases for which required surveillance and control measures are applied and basic biosecurity conditions are met for the purpose of international trade. Such compartments must be clearly documented by the Competent Authority(ies).

COMPETENT AUTHORITY

means the Veterinary Authority or other Governmental Authority of a Member Country having the responsibility and competence for ensuring or supervising the implementation of aquatic animal health and welfare measures, international health certification and other standards and recommendations in the Aquatic Code in the whole territory.

CONTAINER

means a transport appliance:

a) of a permanent type and sufficiently strong to enable repeated use;
b) specially constructed to facilitate transport of aquatic animals or aquatic animal products by one or several means of transport;
c) provided with fittings that make it easy to manipulate, particularly for trans-shipment from one kind of transport vehicle to another;
d) constructed in a watertight way, easy to load and unload and capable of being cleansed and disinfected;
e) ensuring safe and optimal transport of aquatic animals.

CONTINGENCY PLAN

means a documented work plan designed to ensure that all needed actions, requirements and resources are provided in order to eradicate or bring under control outbreaks of specified diseases of aquatic animals.
DIAGNOSIS
means determination of the nature of a disease.

DISEASE
means clinical or non-clinical infection with one or more pathogenic agents.

DISINFECTANTS
means chemical compounds or physical processes capable of destroying pathogenic agents or inhibiting their growth in the course of disinfection.

DISINFECTION
means the process of cleaning and applying disinfectants to inactivate pathogenic agents on potentially contaminated items.

EARLY DETECTION SYSTEM
means an efficient system for ensuring the rapid recognition of signs that are suspicious of a listed disease, or an emerging disease situation, or unexplained mortality, in aquatic animals in an aquaculture establishment or in the wild, and the rapid communication of the event to the Competent Authority, with the aim of activating diagnostic investigation by the Aquatic Animal Health Services with minimal delay. Such a system will include the following characteristics:

a) broad awareness, e.g. among the personnel employed at aquaculture establishments or involved in processing, of the characteristic signs of the listed diseases and emerging diseases;
b) veterinarians or aquatic animal health professionals trained in recognising and reporting suspicions of disease occurrence;
c) ability of the Aquatic Animal Health Services to undertake rapid and effective disease investigation based on a national chain of command;
d) access by the Aquatic Animal Health Services to laboratories with the facilities for diagnosing and differentiating listed diseases and emerging diseases;
e) the legal obligation of private veterinarians or aquatic animal health professionals to report suspicions of disease occurrence to the Competent Authority.

EGG
means a viable fertilised ovum of an aquatic animal. ‘Green eggs’ means newly fertilised ova of fish. ‘Eyed eggs’ means eggs of fish where the eyes of the embryo are visible and that the eggs may be transported.

EMERGING DISEASE
means a disease, other than listed diseases, which has a significant impact on aquatic animal or public health resulting from:

a) a change of known pathogenic agent or its spread to a new geographic area or species; or
b) a newly recognised or suspected pathogenic agent.

EPIDEMIOLOGICAL UNIT
means a group of animals that share approximately the same risk of exposure to a pathogenic agent with a defined location. This may be because they share a common aquatic environment (e.g. fish in a pond, caged fish in a lake), or because management practices make it likely that a pathogenic agent in one group of animals would quickly spread to other animals (e.g. all the ponds on a farm, all the ponds in a village system).

EVISCERATED FISH
means fish from which internal organs, excluding the brain and gills, have been removed.

EXPORTING COUNTRY
means a country from which aquatic animals or aquatic animal products, biological products or pathological material are sent to a destination in another country.

FALLOWING
means, for disease management purposes, an operation where an aquaculture establishment is emptied of aquatic animals susceptible to a disease of concern or known to be capable of transferring the pathogenic agent, and, where feasible, of the carrying water. For aquatic animals of unknown susceptibility and those agreed not to be capable of acting as vectors of a disease of concern, decisions on fallowing should be based on a risk assessment.
**FEED**
means any material (single or multiple), whether processed, semi-processed or raw, as well as live organisms, which is intended to be fed directly to aquatic animals.

**FEED INGREDIENT**
means a component, part or constituent of any combination or mixture making up a feed, including feed additives, whether or not it has a nutritional value in the animal’s diet. Ingredients may be of terrestrial or aquatic, plant or animal origin and may be organic or inorganic substances.

**FREE COMPARTMENT**
means a compartment that fulfils the requirements for self-declaration of freedom from disease with respect to the disease(s) under consideration in accordance with the relevant chapter(s) in the Aquatic Code.

**FREE COUNTRY**
means a country that fulfils the requirements for self-declaration of freedom from disease with respect to the disease(s) under consideration in accordance with the relevant chapter(s) in the Aquatic Code.

**FREE ZONE**
means a zone that fulfils the requirements for self-declaration of freedom from disease with respect to the disease(s) under consideration in accordance with the relevant chapter(s) in the Aquatic Code.

**FRONTIER POST**
means any international airport or any port, railway station or road post open to international trade.

**GAMETES**
means the sperm or unfertilised eggs of aquatic animals that are held or transported separately prior to fertilisation.

**HAZARD**
means a biological, chemical or physical agent in, or a condition of, an aquatic animal or aquatic animal product with the potential to cause an adverse effect on aquatic animal health or public health.

**HEADQUARTERS**
means the Permanent Secretariat of the World Organisation for Animal Health (OIE), located at:
12, rue de Prony, 75017 Paris, FRANCE
Telephone: 33-(0)1 44 15 18 88
Fax: 33-(0)1 42 67 09 87
Electronic mail: oie@oie.int
WWW: http://www.oie.int

**IMPORTING COUNTRY**
means a country that is the final destination to which aquatic animals, aquatic animal products, biological products or pathological material are sent.

**INCIDENCE**
means the number of new outbreaks of disease within a specified period of time in a defined aquatic animal population.

**INFECTED ZONE**
means a zone in which a disease has been diagnosed.

**INFECTION**
means the presence of a multiplying or otherwise developing or latent pathogenic agent in a host. This term is understood to include infestation where the pathogenic agent is a parasite in or on a host.

**INTERNATIONAL AQUATIC ANIMAL HEALTH CERTIFICATE**
means a certificate, issued in conformity with the provisions of Chapter 5.11., describing the aquatic animal health and/or public health requirements that should be fulfilled prior to export of commodity.

**INTERNATIONAL TRADE**
means import, export or transit of aquatic animals, aquatic animal products, biological products and pathological material.
LISTED DISEASES
means diseases that are referred to in Chapter 1.3.

MEAL
means a product derived from an aquatic animal that has been ground and heat processed to reduce the moisture content to less than 10%.

NOTIFICATION
means the procedure by which:
1) the Competent Authority informs the Headquarters,
2) the Headquarters inform Competent Authorities of Member Countries
of the occurrence of a disease in accordance with the provisions of Chapter 1.1.

OUTBREAK
means an occurrence of one or more cases in an epidemiological unit.

PATHOGENIC AGENT
means an organism that causes or contributes to the development of a disease.

PATHOLOGICAL MATERIAL
means samples obtained from live or dead aquatic animals, containing or suspected of containing pathogenic agents, to be sent to a laboratory.

PREVALENCE
means the total number of infected aquatic animals expressed as a percentage of the total number of aquatic animals in a given aquatic animal population at one specific time.

PROBABILITY SAMPLING
means a sampling strategy in which every unit has a known non-zero probability of inclusion in the sample.

PROTECTION ZONE
means a zone established to protect the health status of aquatic animals in a free country or free zone, from those in a country or zone of a different aquatic animal health status, using measures based on the epidemiology of the disease under consideration to prevent spread of the pathogenic agent into a free country or free zone. These measures may include, but are not limited to, vaccination, movement control and an intensified degree of surveillance.

QUARANTINE
means maintaining a group of aquatic animals in isolation with no direct or indirect contact with other aquatic animals, in order to undergo observation for a specified length of time and, if appropriate, testing and treatment, including proper treatment of the effluent waters.

RISK
means the likelihood of the occurrence and the likely magnitude of the biological and economic consequences of an adverse event or effect to animal or human health.

RISK ANALYSIS
means the process composed of hazard identification, risk assessment, risk management and risk communication.

RISK ASSESSMENT
means the scientific evaluation of the likelihood and the biological and economic consequences of entry, establishment and spread of a hazard.

RISK COMMUNICATION
is the interactive exchange of information and opinions throughout the risk analysis process concerning risk, risk-related factors and risk perceptions among risk assessors, risk managers, risk communicators, the general public and other interested parties.
Glossary

RISK MANAGEMENT
means the process of identifying, selecting and implementing measures that can be applied to reduce the level of risk.

SANITARY MEASURE
means a measure, such as those described in various chapters of the Aquatic Code, destined to protect aquatic animal or human health or life within the territory of the Member Country from risks arising from the entry, establishment and/or spread of a hazard.

SELF-DECLARATION OF FREEDOM FROM DISEASE
means declaration by the Competent Authority of the Member Country concerned that the country, zone or compartment is free from a listed disease based on implementation of the provisions of the Aquatic Code and the Aquatic Manual. [NOTE: The Member Country is encouraged to inform the OIE of its claimed status and the OIE may publish the claim but publication does not imply OIE endorsement of the claim.]

SENSITIVITY
means the proportion of true positive tests given in a diagnostic test, i.e. the number of true positive results divided by the number of true positive and false negative results.

SPECIFICITY
means the probability that absence of infection will be correctly identified by a diagnostic test, i.e. the number of true negative results divided by the number of true negative and false positive results.

STAMPING-OUT POLICY
means the carrying out under the authority of the Competent Authority, on confirmation of a disease, of preventive aquatic animal health measures, consisting of killing the aquatic animals that are affected, those suspected of being affected in the population and those in other populations that have been exposed to infection by direct or indirect contact of a kind likely to cause the transmission of the pathogenic agent. All these aquatic animals, vaccinated or unvaccinated, on an infected site should be killed and the carcasses destroyed by burning or burial, or by any other method that will eliminate the spread of infection through the carcasses or products of the aquatic animals destroyed.

This policy should be accompanied by cleansing and disinfection procedures as defined in the Aquatic Code. Fallowing should be for an appropriate period determined by risk assessment.

STUDY POPULATION
means the population from which surveillance data are derived. This may be the same as the target population or a subset of it.

SUBPOPULATION
means a distinct part of a population identifiable in accordance with specific common aquatic animal health characteristics.

SURVEILLANCE
means a systematic series of investigations of a given population of aquatic animals to detect the occurrence of disease for control purposes, and which may involve testing samples of a population.

Susceptible Species
means a species of aquatic animal in which infection has been demonstrated by the occurrence of natural cases or by experimental exposure to the pathogenic agent that mimics natural transmission pathways.

TARGET POPULATION
means, for the purposes of demonstrating freedom from infection, the population of interest, usually made up of all aquatic animals of species susceptible to a specified pathogenic agent in a defined country, zone or aquaculture establishment.

TARGETED SURVEILLANCE
means surveillance targeted at a specific disease or infection.

TERRITORY
means land and water under jurisdiction of a country.
TRANSIT COUNTRY
means a country through which aquatic animals, aquatic animal products, biological products or pathological material destined for an importing country, are transported or in which a stopover is made at a frontier post.

UNIT
means individually identifiable elements. This is a generic concept used to describe, for example, the members of a population, or the elements selected when sampling. In these contexts, examples of units include individual animals, ponds, nets, cages, farms, villages, districts, etc.

VECTOR
means any living organism that transports a pathogenic agent to a susceptible aquatic animal or its food or immediate surroundings. The pathogenic agent may or may not pass through a development cycle within the vector.

VEHICLE
means any method of transport by land, air or water.

VETERINARIAN
means a person with appropriate education, registered or licensed by the relevant veterinary statutory body of a country to practise veterinary medicine/science in that country.

VETERINARY AUTHORITY
means the Governmental Authority of a Member Country, comprising veterinarians, other professionals and paraprofessionals, having the responsibility and competence for ensuring or supervising the implementation of aquatic animal health and welfare measures, international aquatic animal health certification and other standards and recommendations in the Aquatic Code in the whole territory.

VETERINARY STATUTORY BODY
means an autonomous authority regulating veterinarians and veterinary paraprofessionals.

WATER CATCHMENT
means an area or basin of land bounded by natural features such as hills or mountains, into which all run-off water flows.

ZONE
means an area in one or more countries containing an aquatic animal population with a specific aquatic animal health status with respect to a disease, in which surveillance and control measures and basic biosecurity conditions are applied. The zone should be defined by the Competent Authority.

NB: MOST RECENT UPDATE ADOPTED IN 2017.
SECTION 1.

NOTIFICATION, DISEASES LISTED BY THE OIE AND SURVEILLANCE FOR AQUATIC ANIMALS

CHAPTER 1.1.

NOTIFICATION OF DISEASES, AND PROVISION OF EPIDEMIOLOGICAL INFORMATION

Article 1.1.1.

For the purposes of the Aquatic Code and in terms of Articles 5, 9 and 10 of the OIE Organic Statutes, Member Countries shall recognise the right of the Headquarters to communicate directly with the Competent Authority of its territory or territories.

All notifications and all information sent by the OIE to the Competent Authority shall be regarded as having been sent to the country concerned and all notifications and all information sent to the OIE by the Competent Authority shall be regarded as having been sent by the country concerned.

Article 1.1.2.

1) Member Countries shall make available to other Member Countries, through the OIE, whatever information is necessary to minimise the spread of important diseases of aquatic animals and their pathogenic agents and to assist in achieving better worldwide control of these diseases.

2) To achieve this, Member Countries shall comply with the notification requirements specified in Articles 1.1.3. and 1.1.4.

3) For the purposes of this chapter an 'event' means a single outbreak or a group of epidemiologically related outbreaks of a given disease that is the subject of a notification. An event is specific to a pathogen and strain, when appropriate, and includes all related outbreaks reported from the time of the immediate notification through to the final report. Reports of an event include susceptible species, number and geographical distribution of affected aquatic animals and epidemiological units.

4) To assist in the clear and concise exchange of information, reports shall conform as closely as possible to the OIE disease reporting format.

5) The detection of the pathogenic agent of a listed disease in an aquatic animal should be reported, even in the absence of clinical signs. Recognising that scientific knowledge concerning the relationship between pathogenic agents and clinical disease is constantly developing and that the presence of an infectious agent does not necessarily imply the presence of clinical disease, Member Countries shall ensure through their reports that they comply with the spirit and intention of point 1 above.

6) In addition to notifying findings in accordance with Article 1.1.3. and 1.1.4., Member Countries shall also provide information on the measures taken to prevent the spread of diseases. Information shall include quarantine measures and restrictions applied to the movement of aquatic animals, aquatic animal products, biological products and other miscellaneous objects which could by their nature be responsible for transmission of disease. In the case of diseases transmitted by vectors, the measures taken against such vectors shall also be specified.
Article 1.1.3.

The Competent Authority shall, under the responsibility of the Delegate, send to the Headquarters:

1) in accordance with relevant provisions in the disease-specific chapters, notification, through the World Animal Health Information System (WAHIS) or by fax or e-mail within 24 hours of any of the following events:
   a) first occurrence of a listed disease in a country, a zone or a compartment;
   b) recurrence of a listed disease in a country, a zone or a compartment following the final report that declared the outbreak ended;
   c) first occurrence of a new strain of a pathogenic agent of a listed disease in a country, a zone or a compartment;
   d) a sudden and unexpected change in the distribution or increase in incidence or virulence of, or morbidity or mortality caused by the pathogenic agent of a listed disease, present within a country, a zone or a compartment;
   e) occurrence of a listed disease in a new host species;

2) weekly reports subsequent to a notification under point 1 above, to provide further information on the evolution of the event which justified the notification. These reports should continue until the disease has been eradicated or the situation has become sufficiently stable so that six-monthly reporting under point 3 will satisfy the obligation of the Member Country. For each event notified, a final report should be submitted;

3) six-monthly reports on the absence or presence and evolution of listed diseases and information of epidemiological significance to other Member Countries;

4) annual reports concerning any other information of significance to other Member Countries.

Article 1.1.4.

Competent Authorities shall, under the responsibility of the Delegate, send to the Headquarters:

1) a notification through WAHIS or by fax or email, when an emerging disease event has occurred in a country, a zone or a compartment;

2) periodic reports subsequent to a notification of an emerging disease should continue:
   a) for the time necessary to have reasonable certainty that:
      i) the disease has been eradicated; or
      ii) the situation has become stable;
   OR
   b) until sufficient scientific information is available to determine whether it meets the criteria for inclusion in the OIE list as described in Chapter 1.2.;

3) a final report once requirements in point 2a) or b) are met.

Article 1.1.5.

1) The Competent Authority of a country in which an infected zone or compartment is located shall inform the Headquarters when this country, zone or compartment becomes free from the disease.

2) A country, zone or compartment may be considered to have regained freedom from a specific disease when all relevant conditions given in the Aquatic Code have been fulfilled.

3) The Competent Authority of a Member Country which establishes one or several free zones or free compartments shall inform the Headquarters, giving necessary details, including the criteria on which the free status is based, the requirements for maintaining the status and indicating clearly the location of the zones or compartments on a map of the territory of the Member Country.

Article 1.1.6.

1) Although Member Countries are only required to notify listed diseases, and emerging diseases, they are encouraged to provide the OIE with other important aquatic animal health information.
2) The Headquarters shall communicate by email or through the interface of WAHIS to Competent Authorities all notifications received as provided in Articles 1.1.2. to 1.1.5. and other relevant information.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2016.
CHAPTER 1.2.

CRITERIA FOR LISTING AQUATIC ANIMAL DISEASES

Article 1.2.1.

Introduction

This chapter describes the criteria for the inclusion of diseases in Chapter 1.3.

The objective of listing diseases is to support Member Countries by providing information needed to take appropriate action to prevent the transboundary spread of important diseases of aquatic animals. This is achieved by transparent, timely and consistent notification.

Each listed disease usually has a corresponding chapter that assists Member Countries in the harmonisation of disease detection, prevention and control, and provides standards for safe international trade in aquatic animals and aquatic animal products.

The requirements for notification are detailed in Chapter 1.1.

Principles and methods of validation of diagnostic tests are described in Chapter 1.1.2. of the Aquatic Manual.

Article 1.2.2.

The criteria for the inclusion of a disease in the OIE list are as follows:

1) International spread of the pathogenic agent (via aquatic animals, aquatic animal products, vectors or fomites) is likely.

AND

2) At least one country may demonstrate country or zone freedom from the disease in susceptible aquatic animals, based on provisions of Chapter 1.4.

AND

3) A precise case definition is available and a reliable means of detection and diagnosis exists.

AND

4) a) Natural transmission to humans has been proven, and human infection is associated with severe consequences.

OR

b) The disease has been shown to affect the health of cultured aquatic animals at the level of a country or a zone resulting in significant consequences e.g. production losses, morbidity or mortality at a zone or country level.

OR

c) The disease has been shown to, or scientific evidence indicates that it would affect the health of wild aquatic animals resulting in significant consequences e.g. morbidity or mortality at a population level, reduced productivity or ecological impacts.

NB: FIRST ADOPTED IN 2003; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 1.3.

DISEASES LISTED BY THE OIE

Preamble: The following diseases are listed by the OIE in accordance with the criteria for listing an aquatic animal disease (see Article 1.2.2.).

In case of modifications of this list of aquatic animal diseases adopted by the World Assembly of Delegates, the new list comes into force on 1 January of the following year.

Article 1.3.1.

The following diseases of fish are listed by the OIE:
- Epizootic haematopoietic necrosis disease
- Infection with Aphanomyces invadans (epizootic ulcerative syndrome)
- Infection with Gyrodactylus salaris
- Infection with HPR-deleted or HPR0 infectious salmon anaemia virus
- Infection with salmonid alphavirus
- Infectious haematopoietic necrosis
- Koi herpesvirus disease
- Red sea bream iridoviral disease
- Spring viraemia of carp
- Viral haemorrhagic septicaemia.

Article 1.3.2.

The following diseases of molluscs are listed by the OIE:
- Infection with abalone herpesvirus
- Infection with Bonamia ostreae
- Infection with Bonamia exitiosa
- Infection with Marteilia refringens
- Infection with Perkinsus marinus
- Infection with Perkinsus olseni
- Infection with Xenohaliotis californiensis.

Article 1.3.3.

The following diseases of crustaceans are listed by the OIE:
- Acute hepatopancreatic necrosis disease
- Infection with Aphanomyces astaci (crayfish plague)
- Infection with Hepatobacter penaei (necrotising hepatopancreatitis)
- Infection with infectious hypodermal and haematopoietic necrosis virus
- Infection with infectious myonecrosis virus
- Infection with Macrobrachium rosenbergii nodavirus (white tail disease)
- Infection with Taura syndrome virus
- Infection with white spot syndrome virus
- Infection with yellow head virus genotype 1.
Article 1.3.4.

The following diseases of amphibians are listed by the OIE:
- Infection with *Batrachochytrium dendrobatidis*
- Infection with *Batrachochytrium salamandrivorans*
- Infection with *Ranavirus* species

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 1.4.

AQUATIC ANIMAL HEALTH SURVEILLANCE

Article 1.4.1.

Introduction and objectives

1) Surveillance activities may be performed to achieve any of the following objectives:
   a) demonstrating the absence of disease;
   b) identifying events requiring notification as listed in Article 1.1.3.;
   c) determining the occurrence or distribution of endemic disease, including changes to their incidence or prevalence (or its contributing factors), in order to:
      i) provide information for domestic disease control programmes,
      ii) provide relevant disease occurrence information to be used by trading partners for qualitative and quantitative risk assessment.

   The type of surveillance applied depends on the desired outputs needed to support decision-making. Surveillance data determine the quality of disease status reports and should satisfy information requirements for accurate risk analysis both for international trade as well as for national decision-making. Surveillance of endemic diseases provides valuable information for day-to-day health management and can act as the foundation for detecting outbreaks of exotic disease and demonstrating specific disease freedom.

   Surveillance systems described in this chapter should also be used to generate information for decisions on prescribed disease prevention and control programmes. However, the actual strategies for prevention and control are beyond the scope of this chapter on surveillance recommendations.

   Having a suitable management strategy to respond to surveillance data is of utmost importance for the successful implementation of surveillance systems.

2) Essential prerequisites to enable a Member Country to provide information for the evaluation of its animal health status are:
   a) that the particular Member Country complies with the provisions of Chapter 3.1 on the quality of the Aquatic Animal Health Services;
   b) that, where possible, surveillance data be complemented by other sources of information (e.g. scientific publications, research data, documented field observations and other non-survey data);
   c) that transparency in the planning and execution of surveillance activities and the analysis and availability of data and information, be maintained at all times, in accordance with Chapter 1.1.

3) The following recommendations may be applied to all diseases, their agents, and susceptible species as listed in the Aquatic Manual, and are designed to assist with the development of surveillance methodologies. Where possible, the development of surveillance systems using these recommendations should be based on the relevant information in the individual disease chapters in the Aquatic Manual. These recommendations are also applicable to non-listed diseases that may be of importance to a country or region, such as new or emerging diseases. There is sometimes a perception that surveillance can only be conducted using sophisticated methodologies. However, an effective surveillance system can also be developed by making use of gross observations and already available resources.

   More detailed information in each disease chapter (where it exists) of the Aquatic Manual may be used to further refine the general approaches described in this chapter. Where detailed disease-specific information is not available, surveillance can also be conducted following the recommendations in this chapter. Access to epidemiological expertise would be invaluable for the design, implementation of the system and interpretation of results derived from a surveillance system.
Chapter 1.4.- Aquatic animal health surveillance

Article 1.4.2.

Principles of surveillance

1) Surveillance may be based on many different data sources and can be classified in a number of ways, including:
   a) the means by which data are collected (targeted versus non-targeted);
   b) the disease focus (pathogen-specific versus general surveillance); and
   c) the way in which units for observation are selected (surveys versus non-random data sources).

2) Surveillance activities include:
   a) population-based surveys, such as:
      i) systematic sampling at slaughter;
      ii) random surveys;
   b) non-random surveillance activities, such as:
      i) disease reporting or notifications;
      ii) control programmes/health schemes;
      iii) targeted testing/screening;
      iv) post-mortem inspections;
      v) laboratory investigation records;
      vi) biological specimen banks;
      vii) sentinel units;
      viii) field observations;
      ix) farm production records.

3) In addition, surveillance data should be supported by related information, such as:
   a) data on the epidemiology of the disease, including environmental, and host and wild reservoir population distributions;
   b) data on farmed and wild animal movements and trading patterns for aquatic animals and aquatic animal products, including potential for exposure to populations of wild aquatic animals, water sources or other contacts;
   c) national animal health regulations, including information on compliance with them and their effectiveness;
   d) history of imports of potentially infected material; and
   e) biosecurity measures in place.

4) The sources of evidence should be fully described. A survey should include a description of the sampling strategy used for the selection of units for testing. For non-random data sources, a full description of the system is required including the source(s) of the data, when the data were collected, and a consideration of any biases that may be inherent in the system.

Article 1.4.3.

Critical elements of surveillance

In assessing the quality of a surveillance system, the following critical elements need to be addressed.

1. Populations
   Ideally, surveillance should be carried out in such a way as to take into account all animal species susceptible to the disease in a country, zone or compartment. The surveillance activity may cover all individuals in the population or part of them. Estimates of total population at risk for each species are required. When surveillance is conducted only on a subpopulation, care should be taken regarding the inferences made from the results.

   For listed diseases, definitions of appropriate populations should be based on the specific recommendations of the disease chapters of the Aquatic Manual.

2. Epidemiological unit
   The relevant epidemiological unit for the surveillance system should be defined and documented to ensure that it is representative of the population or targeted subpopulations that would generate the most useful inferences about disease patterns. Therefore, it should be chosen taking into account factors such as reservoirs, vectors, immune status, genetic resistance and age, sex, and other host criteria.
3. Clustering

*Disease* in a country, zone or *compartment* usually clusters rather than being uniformly or randomly distributed through a population. Clustering of *disease* may occur in space (e.g. tank, pond, farm, or *compartment*), time (e.g. season), or animal subgroups (e.g. age, physiological condition). Clustering should be taken into account in the design of *surveillance* activities and interpretation of *surveillance* data.

4. Case and outbreak definitions

Clear and unambiguous *case definitions* and outbreak definitions should be developed and documented for each *disease* under *surveillance*, using, where they exist, the standards in this chapter and the *Aquatic Manual*.

5. Analytical methodologies

*Surveillance* data should be analysed using appropriate methodologies, and at the appropriate organisational levels to facilitate effective decision-making, whether it be planning interventions or demonstrating status. Methodologies for the analysis of *surveillance* data should be flexible to deal with the complexity of real life situations. No single method is applicable in all cases. Different methodologies may be needed to accommodate the relevant pathogens, varying production and *surveillance* systems, and types, quality, and amounts of data/information available.

The methodology used should be based on the best available information that is in accord with current scientific thinking. The methodology should be in accordance with this chapter and fully documented, and supported by reference to the scientific literature and other sources, including expert opinion. Sophisticated mathematical or statistical analyses should only be carried out when justified by the proper amount and quality of field data.

Consistency in the application of different methodologies should be encouraged and transparency is essential in order to ensure fairness and rationality, consistency in decision-making and ease of understanding. The uncertainties, assumptions made, and the effect of these on the final conclusions should be documented.

6. Testing

*Surveillance* involves the detection of *disease* by the use of appropriate *case definitions* based on the results of one or more tests for evidence of *disease* status. In this context, a test may range from detailed laboratory examinations to field observations and the analysis of production records. The performance of a test at the population level (including field observations) may be described in terms of its *sensitivity* and *specificity* and predictive values. Imperfect *sensitivity* and/or *specificity* will have an impact on the conclusions from *surveillance*. Therefore, these parameters should be taken into account in the design of *surveillance* systems and analysis of *surveillance* data as described in this chapter.

Although not determined for many *aquatic animal* diseases, *sensitivity* and *specificity* should be estimated as best as possible for a specific testing situation. Alternatively, where values for *sensitivity* and/or *specificity* for a particular test and testing situation are estimated in the *disease* chapter in the *Aquatic Manual*, these values may be used as a guide.

Samples from a number of *aquatic animals* or units may be pooled and subjected to a testing protocol. The results should be interpreted using *sensitivity* and *specificity* values that have been determined or estimated for that particular pool size and testing procedure.

7. Quality assurance

*Surveillance* systems should incorporate the principles of quality assurance and be subjected to periodic auditing to ensure that all components of the system function and provide verifiable documentation of procedures and basic checks to detect significant deviations of procedures from those documented in the design.

8. Validation

Results from animal health *surveillance* systems are subject to one or more potential *biases*. When assessing the results, care should be taken to identify potential *biases* that can inadvertently lead to an over-estimate or an under-estimate of the parameters of interest.

9. Data collection and management

The success of a *surveillance* system is dependent on a reliable process for data collection and management. The process may be based on paper records or computerised. Even where data are collected for non-survey purposes (e.g. during *disease* control interventions, inspections for movement control or during *disease* eradication schemes), the consistency and quality of data collection and event reporting in a format that facilitates analysis, is critical. Factors influencing the quality of collected data include:

a) the distribution of, and communication between, those involved in generating and transferring data from the field to a centralised location;
b) motivation of the people involved in the surveillance system;

c) the ability of the data processing system to detect missing, inconsistent or inaccurate data, and to address these problems;

d) maintenance of disaggregated data rather than the compilation of summary data;

e) minimisation of transcription errors during data processing and communication.

Article 1.4.4.

Population-based surveys

In addition to the principles for surveillance discussed in Article 1.4.6., the following recommendations should be used when planning, implementing and analysing surveys.

1. Types of surveys

Surveys may be conducted on the entire target population (i.e., a census) or on a sample. Periodic or repeated surveys conducted in order to document disease freedom should be done using probability based sampling methods (simple random selection, cluster sampling, stratified sampling, systematic sampling) so that data from the study population can be extrapolated to the target population in a statistically valid manner. Non-probability based sampling methods (convenience, expert choice, quota) can also be used. Recognising the inherent impracticalities in sampling from some aquatic animal populations, non-probability based sampling could be used when biases are recognised and used to optimise detection.

The sources of information should be fully described and should include a detailed description of the sampling strategy used for the selection of units for testing. Also, consideration should be made of any biases that may be inherent in the survey design.

2. Survey design

The population of epidemiological units should first be clearly defined; hereafter sampling units appropriate for each stage, depending on the design of the survey, should be defined.

The design of the survey will depend on the size and structure of the population being studied, the epidemiology of the disease and the resources available.

3. Sampling

The objective of sampling from a population is to select a subset of units from the population that is representative of the population with respect to the object of the study such as the presence or absence of disease. Sampling should be carried out in such a way as to provide the best likelihood that the sample will be representative of the population, within the practical constraints imposed by different environments and production systems. In order to detect the presence of a disease in a population of unknown disease status, sampling methods that optimise the detection of disease can be used. In such cases, care should be taken regarding the inferences made from the results.

4. Sampling methods

When selecting epidemiological units from within a population the objectives of the surveillance system should be considered. In general, probability sampling (e.g., simple random selection) is preferable. When this is not possible, sampling should provide the best practical chance of generating optimal inferences about disease patterns in the target population.

In any case, the sampling method used at all stages should be fully documented and justified.

5. Sample size

In general, surveys are conducted either to demonstrate the presence or absence of a factor (e.g., disease) or to estimate a parameter (e.g., the prevalence of disease). The method used to calculate sample size for surveys depends on the purpose of the survey, the expected prevalence (also referred to as the threshold prevalence), the level of confidence desired of the survey results and the performance (e.g., sensitivity and specificity estimates) of the tests used.
Non-random data sources used in surveillance

Surveillance systems routinely use non-random data, either alone or in combination with surveys.

1. Common non-random surveillance data sources

A wide variety of non-random surveillance data sources may be available. These vary in their primary purpose and the type of surveillance information they are able to provide. Some surveillance systems are primarily established as early detection systems, but may also provide valuable information to demonstrate freedom from disease. Other systems provide cross-sectional information suitable for prevalence estimation, either once or repeatedly, while yet others provide continuous information, suitable for the estimate of incidence data (e.g. disease reporting systems, sentinel sites, testing schemes).

a) Disease reporting or notification system

Data derived from disease reporting systems can be used in combination with other data sources to substantiate claims of animal health status, to generate data for risk analysis, or for early detection. The first step of a disease reporting or notification system is often based on the observation of abnormalities (e.g. clinical signs, reduced growth, elevated mortality rates, behavioural changes, etc.), which can provide important information about the occurrence of endemic, exotic or new diseases. Effective laboratory support is, however, an important component of most reporting systems. Reporting systems relying on laboratory confirmation of suspect clinical cases should use tests that have a high specificity. Reports should be released by the laboratory in a timely manner, with the amount of time from disease detection to report generation minimised.

b) Control programmes/health schemes

Animal disease control programmes or health schemes, while focusing on the control or eradication of specific diseases, should be planned and structured in such a manner as to generate data that are scientifically verifiable and contribute to surveillance.

c) Targeted sampling

This may involve sampling targeted to selected sections of the population (subpopulations), in which disease is more likely to be introduced or found. Examples include selecting culled and dead animals for testing, animals exhibiting clinical signs, animals located in a defined geographical area and specific age or commodity group.

d) Post-harvest inspections

Inspections of aquatic animal slaughter premises or processing plants may provide valuable surveillance data provided diseased aquatic animals survive to slaughter. Post-harvest inspections are likely to provide good coverage only for particular age groups and geographical areas. Post-harvest surveillance data are subject to obvious biases in relation to target population and study population (e.g. only animals of a particular class and age may be slaughtered for human consumption in significant numbers). Such biases need to be recognised when analysing surveillance data.

Both for traceback in the event of detection of disease and for analysis of spatial and population-level coverage, there should be, if possible, an effective identification system that relates each animal in the slaughter premises/processing plant to its locality of origin.

e) Laboratory investigation records

Analysis of laboratory investigation records may provide useful surveillance information. The coverage of the system will be increased if analysis is able to incorporate records from national, accredited, university and private sector laboratories. Valid analysis of data from different laboratories depends on the existence of standardised diagnostic procedures and standardised methods for interpretation and data recording. If available, the method listed in the Aquatic Manual in relation to the purpose of testing should be used. As with post-harvest inspections, there needs to be a mechanism to relate specimens to the farm of origin. It should be recognised that laboratory submissions may not accurately reflect the disease situation on the farm.

f) Biological specimen banks

Specimen banks consist of stored specimens, gathered either through representative sampling or opportunistic collection or both. Specimen banks may contribute to retrospective studies, including providing support for claims of historical freedom from disease, and may allow certain studies to be conducted more quickly and at lower cost than alternative approaches.

g) Sentinel units

Sentinel units/sites involve the identification and regular testing of one or more of animals of known health/exposure status in a specified geographical location to detect the occurrence of disease. They are particularly useful for surveillance of diseases with a strong spatial component, such as vector borne diseases. Sentinel units provide the opportunity to target surveillance depending on the likelihood of disease...
(related to vector habitats and host population distribution), cost and other practical constraints. Sentinel units may provide evidence of freedom from disease, or provide data on prevalence and incidence as well as the distribution of disease. Cohabitation of sentinel units (preferably of the most susceptible species and life stage) with a susceptible population should be considered for testing disease in populations of valuable animals, the lethal sampling of which may be unacceptable (e.g. ornamental fish) or in animal subpopulations where sampling techniques are incapable of detecting the presence of disease or infection (e.g. where vaccination means that serological tests are inapplicable).

h) Field observations

Clinical observations of epidemiological units in the field are an important source of surveillance data. The sensitivity and/or specificity of field observations may be relatively low, but these can be more easily determined and controlled if a clear, unambiguous and easy to apply standardised case definition is applied. Education of potential field observers in application of the case definition and reporting is an important component. Ideally, both the number of positive observations and the total number of observations should be recorded.

i) Farm production records

Systematic analysis of farm production records may be used as an indicator of the presence or absence of disease at the population level. If production records are accurate and consistently maintained, the sensitivity of this approach may be quite high (depending on the disease), but the specificity is often quite low.

2. Critical elements for non-random data used in surveillance

There are a number of critical factors that should be taken into account when using non-random surveillance data such as coverage of the population, duplication of data, and sensitivity and specificity of tests that may give rise to difficulties in the interpretation of data. Surveillance data from non-random data sources may increase the level of confidence or be able to detect a lower level of prevalence with the same level of confidence compared to surveys.

3. Analytical methodologies

Different scientifically valid methodologies may be used for the analysis of non-random surveillance data. This most often requires information on parameters of importance to the surveillance system, such as sensitivity and specificity and prior probabilities of infection, i.e. apparent prevalences (e.g. for predictive value calculations). Where no such data are available, estimates based on expert opinions, gathered and combined using a formal, documented and scientifically valid methodology may be used.

4. Combination of multiple sources of data

The methodology used to combine the evidence from multiple or recurrent (e.g. time series) data sources should be scientifically valid, and fully documented including references to published material.

Surveillance information gathered from the same country, zone or compartment at different times (e.g. repeated annual surveys) may provide cumulative evidence of animal health status. Such evidence gathered over time may be combined to provide an overall level of confidence. However, a single larger survey, or the combination of data collected during the same time period from multiple random or non-random sources, may be able to achieve the same level of confidence in a shorter period of time.

Analysis of surveillance information gathered intermittently or continuously over time should, where possible, incorporate the time of collection of the information to take into account the decreased value of older information. The sensitivity, specificity and completeness of data from each source should also be taken into account for the final overall confidence level estimation.

Article 1.4.6.

Pathways to demonstrate freedom from disease

The different paths to declaration of freedom from disease are summarised in the diagram below.
1. **Absence of susceptible species**

   Unless otherwise specified in the relevant disease chapter, a country, zone or compartment may be recognised as being free from disease without applying targeted surveillance if there are no susceptible species (as listed in the relevant chapter of this Aquatic Manual, or in the scientific literature) present in that country, zone or compartment.

2. **Historically free**

   Unless otherwise specified in the relevant disease chapter, a country, zone or compartment may be declared free from disease without formally applying a pathogen-specific surveillance programme when:
   
   a) there has never been a substantiated occurrence of disease reported officially or in the scientific literature (peer reviewed), or
   
   b) disease has not occurred for at least ten years, provided that the pathogenic agents are likely to produce identifiable clinical signs in observable susceptible animals,

   and for at least the past ten years:
   
   c) the basic biosecurity conditions are in place and effectively enforced;
   
   d) no vaccination against the disease has been carried out unless otherwise allowed for in the Aquatic Code;
   
   e) disease is not known to be established in wild aquatic animals within the country or zone intended to be declared free. (A country or zone cannot apply for historical freedom if there is any evidence of disease in wild aquatic animals. However, specific surveillance in wild aquatic animals is not necessary.)

   A country, zone or compartment that was self-declared free on the basis of the absence of susceptible species, but subsequently introduces any of the susceptible species as listed in the Aquatic Manual, may be considered historically free from the disease provided that:
   
   f) the country, zone or compartment of origin was declared free of the disease at the time of introduction;
   
   g) basic biosecurity conditions were introduced prior to the introduction;
   
   h) no vaccination against the disease has been carried out unless otherwise allowed for in the disease-specific chapter of this Aquatic Code.

3. **Last occurrence within the previous ten years/previously unknown status**

   Countries, zones or compartments that have achieved eradication (or in which the disease has ceased to occur) within the previous ten years or where the disease status is unknown, should follow the pathogen-specific
surveillance requirements in the Aquatic Manual if they exist. In the absence of disease-specific information to aid the development of a surveillance system, declaration of disease freedom should follow at least two surveys per year (for at least two consecutive years) to be conducted three or more months apart, on the appropriate species, at the appropriate life stage and at times of the year when temperature and season offer the best opportunity to detect the pathogen. Surveys should be designed to provide an overall 95% confidence or greater and with a design prevalence at the animal and higher levels of aggregation (i.e. pond, farm, village, etc.) of 2% or lower (this value may be different for different diseases and may be provided in the disease-specific chapter in the Aquatic Manual). Such surveys should not be based on voluntary submission and should be developed following the recommendations provided in the Aquatic Manual. Survey results will provide sufficient evidence of disease freedom provided that for at least the past ten years these additional criteria are met:

a) the basic biosecurity conditions are in place and effectively enforced;

b) no vaccination against the disease has been carried out unless otherwise provided in the Aquatic Code;

c) disease is not known to be established in wild aquatic animals within the country or zone intended to be declared free. (A country or zone cannot apply for freedom if there is any evidence of disease in wild aquatic animals. Specific surveillance in wild aquatic animals of susceptible species is necessary to confirm absence.)

Article 1.4.7.

Maintenance of disease free status

A country or zone that has been declared free from disease following the provisions of the Aquatic Code may discontinue pathogen-specific surveillance while maintaining the disease free status provided that:

1) if present, the pathogen is likely to produce identifiable clinical signs in observable susceptible species;

2) the basic biosecurity conditions are in place and effectively enforced;

3) no vaccination against the disease has been carried out unless otherwise provided in the Aquatic Code;

4) where applicable, surveillance has previously demonstrated that disease is not present in populations of wild aquatic animal of susceptible species.

A special case can be made for a disease free compartment in a country or zone not declared disease free, surveillance should be maintained at a level commensurate with the degree of risk and exposure to potential sources of disease is prevented.

Article 1.4.8.

Design of surveillance programmes to demonstrate freedom from disease

A surveillance programme to demonstrate freedom from disease should meet the following requirements in addition to the general requirements for surveillance outlined in this chapter.

Freedom from disease implies the absence of the pathogenic agent in the country, zone or compartment. Scientific methods cannot provide absolute certainty of the absence of disease. Demonstrating freedom from disease involves providing sufficient evidence to demonstrate (to a level of confidence acceptable to Member Countries) that disease with a specified pathogen is not present in a population. In practice, it is not possible to prove (i.e. be 100% confident) that a population is free from disease. Instead, the aim is to provide adequate evidence (to an acceptable level of confidence), that disease, if present, is present in less than a specified proportion of the population (i.e. threshold prevalence).

However, apparent disease at any level in the target population automatically invalidates any freedom from disease claim unless the positive test results are accepted as false positives based on specificity values described in the relevant disease chapter.

The provisions of this Article are based on the principles described above and the following premises:

- in the absence of disease and vaccination, the farmed and wild animal populations would become susceptible over a period of time;
- the pathogenic agents to which these provisions apply are likely to produce identifiable clinical signs in observable susceptible animals;
- to increase the probability of detecting the specific pathogenic agent, the susceptibility of the aquatic animal and the timing of sampling should be under appropriate conditions;
- the Aquatic Animal Health Services will be able to investigate, diagnose and report disease, if present;
– the appropriate diagnostic method as described in the Aquatic Manual be used;
– any claim for the absence of disease over a long period of time in a susceptible population can be substantiated by effective disease investigation and reporting by a Member Country.

1. Objectives

The objective of this kind of surveillance system is to contribute on an on-going basis evidence to demonstrate freedom from disease in a particular country, zone or compartment with a known confidence and reference to a predetermined design prevalence and diagnostic test characteristics. The level of confidence and the design prevalence will depend on the testing situation, disease and host population characteristics and on the resources available.

A single such survey can contribute evidence adding to an on-going collection of health data. However, single surveys in isolation rarely, if ever, provide sufficient evidence that an aquatic animal disease is absent and should be augmented with on-going targeted evidence collection (e.g. ongoing disease sampling or passive detection capabilities) to substantiate claims of freedom from disease.

2. Population

The population of epidemiological units should be clearly defined. The target population consists of all individuals of all susceptible species to the disease in a country, zone or compartment to which the surveillance results apply. Sometimes components of the target population are at higher risk of being the point of introduction for an exotic disease. In these cases, it is advisable to focus surveillance efforts on this part of the population, such as farms on a geographical border.

The design of the survey will depend on the size and structure of the population being studied. If the population is relatively small and can be considered to be homogenous with regards to risk of infection, a single-stage survey can be used. If different of the same aquaculture establishment do not share water, they may be considered as epidemiologically separate populations.

In larger populations where a sampling frame is not available, or when there is a likelihood of clustering of disease, multi-stage sampling is required. In two-stage sampling, at the first stage of sampling, groups of animals (e.g. ponds, farms or villages) are selected. At the second stage, animals are selected for testing from each of the selected groups.

In the case of a complex (e.g. multi-level) population structure, multi-level sampling may be used and the data analysed accordingly.

3. Sources of evidence

Surveillance data may originate from a number of different sources, including:

a) population-based surveys using one or more tests to detect the aetiological agent or evidence of infection;
b) other non-random sources of data, such as:
   i) sentinel sites;
   ii) disease notifications and laboratory investigation records;
   iii) academic and other scientific studies;
c) a knowledge of the biology of the agent, including environmental, host population distribution, known geographical distribution, vector distribution and climatic information;
d) history of imports of potentially infected material;
e) biosecurity measures in place;
f) any other sources of information that provide contributory evidence regarding disease in the country, zone or compartment.

The sources of evidence should be fully described. A survey should include a description of the sampling strategy used for the selection of units for testing. For complex surveillance systems, a full description of the system is required including consideration of any biases that may be inherent in the system. Evidence to support claims of freedom from disease can use non-random sources of information provided that, overall, any biases introduced subsequently favour the detection.
4. **Statistical methodology**

Analysis of test results from a survey shall be in accordance with the provisions of this chapter and consider the following factors:

a) the survey design;

b) the sensitivity and specificity of the test, or test system;

c) the design prevalence (or prevalences where a multi-stage design is used);

d) the results of the survey.

Analysis of data for evidence of freedom from infection involves estimating the probability (alpha) that the evidence observed (the results of surveillance) could have been produced under the null hypothesis that infection is present in the population at a specified prevalence(s) (the design prevalences). The confidence in (or, equivalently, the sensitivity of) the surveillance system that produced the evidence is equal to 1–alpha. If the confidence level exceeds a pre-set threshold, the evidence is deemed adequate to demonstrate freedom from infection.

The required level of confidence in the surveillance system (probability that the system would detect infection if infection were present at the specified level) should be greater than or equal to 95%.

The power (probability that the system would report that no infection is present if infection is truly not present) may be set to any value. By convention, this is often set to 80%, but may be adjusted in accordance with the country’s or zone’s requirements.

Different statistical methodologies for the calculation of the probability alpha, including both quantitative and qualitative approaches, are acceptable as long as they are based on accepted scientific principles.

The methodology used to calculate the confidence in the surveillance system should be scientifically based and clearly documented, including references to published work describing the methodology.

Statistical analysis of surveillance data often requires assumptions about population parameters or test characteristics. These are usually based on expert opinion, previous studies on the same or different populations, expected biology of the agent, and so on. The uncertainty around these assumptions should be quantified and considered in the analysis (e.g. in the form of prior probability distributions in a Bayesian setting).

For surveillance systems used to demonstrate freedom from specific diseases, calculation of the confidence of a surveillance system is based on the null hypothesis that infection is present in the population. The level of infection is specified by the design prevalence. In the simplest case, this is the prevalence of infection in a homogenous population. More commonly, in the presence of a complex (e.g. multi-level) population structure more than one design prevalence value is required, for instance, the animal-level prevalence (proportion of infected animals in an infected farm) and the group-level prevalence (proportion of infected farms in the country, zone or compartment).

Further levels of clustering may be considered, requiring further design prevalence values.

The values for design prevalence used in calculations should be those specified in the relevant disease chapter (if present) of the Aquatic Manual. If not specified for the particular disease, justification for the selection of design prevalence values should be provided, and should be based on the following recommendations:

- At the individual animal level, the design prevalence is based on the biology of the infection in the population. It is equal to the minimum expected prevalence of infection in the study population, if the infection had become established in that population. It is dependent on the dynamics of infection in the population and the definition of the study population (which may be defined to maximise the expected prevalence in the presence of infection).

- A suitable design prevalence value at the animal level (e.g. prevalence of infected animals in a cage) may be:

  - between 1% and 5% for infections that are present in a small part of the population e.g. are transmitted slowly or are at the early stages of an outbreak of disease, etc.;
  
  - over 5% for highly transmissible infections.

If reliable information, including expert opinion, on the expected prevalence in an infected population is not available, a value of 2% should be used for the design prevalence.

- At higher levels (e.g. cage, pond, farm, village, etc.) the design prevalence usually reflects the prevalence of infection that is practically and reasonably able to be detected by a surveillance system. Detection of infection at the lowest limit (a single infected unit in the population) is rarely feasible in large populations. The expected
behaviour of the *infection* may also play a role. *Infections* that have the ability to spread rapidly between farms may have a higher farm-level design *prevalence* than slow-moving *infections*.

A suitable design *prevalence* value for the first level of clustering (e.g. proportion of infected farms in a zone) is normally not greater than 2%. If a higher design *prevalence* is selected, it should be justified.

When *surveillance* data are used to estimate *incidence* and *prevalence* measures for the purpose of describing disease occurrence in terms of animal unit, time and place, these measures can be calculated for an entire population and specific time period, or for subsets defined by host characteristics (e.g. age-specific *incidence*). *Incidence* estimation requires on-going *surveillance* to detect new cases while *prevalence* is the estimated proportion of infected individuals in a population at a given time point. The estimation process should consider test sensitivity and specificity.

5. **Clustering of infection**

*Infection* in a country, zone or compartment usually clusters rather than being uniformly distributed through a population. Clustering may occur at a number of different levels (e.g. a cluster of moribund fish in a pond, a cluster of ponds in a farm, or a cluster of farms in a zone). Except when dealing with demonstrably homogenous populations, *surveillance* should take this clustering into account in the design and the statistical analysis of the data, at least at what is judged to be the most significant level of clustering for the particular animal population and *infection*.

6. **Test characteristics**

All *surveillance* involves performing one or more tests for evidence of the presence of current or past *infection*, ranging from detailed laboratory examinations to farmer observations. The performance level of a test at the population level is described in terms of its sensitivity and specificity. Imperfect sensitivity and/or specificity impact on the interpretation of *surveillance* results and should be taken into account in the analysis of *surveillance* data. For example, in the case of a test with imperfect specificity, if the population is free of disease or has a very low *prevalence* of *infection*, all or a large proportion of positive tests will be false. Subsequently, samples that test positive can be confirmed or refuted using a highly specific test. Where more than one test is used in a *surveillance* system (sometimes called using tests in series or parallel), the sensitivity and specificity of the test combination should be calculated.

All calculations should take the performance level (sensitivity and specificity) of any tests used into account. The values of sensitivity and specificity used for calculations should be specified, and the method used to determine or estimate these values should be documented. Test sensitivity and specificity can be different when applied to different populations and testing scenarios. For example, test sensitivity may be lower when testing carrier animals with low level *infections* compared to moribund animals with clinical disease. Alternatively, specificity depends on the presence of cross-reacting agents, the distribution of which may be different under different conditions or regions. Ideally, test performance should be assessed under the conditions of use otherwise increased uncertainty exists regarding their performance. In the absence of local assessment of tests, values for sensitivity and/or specificity for a particular test that are specified in the *Aquatic Manual* may be used but the increased uncertainty associated with these estimates should be incorporated into the analysis of results.

Pooled testing involves the pooling of specimens from multiple individuals and performing a single test on the pool. Pooled testing is an acceptable approach in many situations. Where pooled testing is used, the results of testing should be interpreted using sensitivity and specificity values that have been determined or estimated for that particular pooled testing procedure and for the applicable pool sizes being used. Analysis of the results of pooled testing should, where possible, be performed using accepted, statistically based methodologies, which should be fully documented, including published references.

When applied to a *surveillance* system, the probabilities of correct assessment of the health status of the *epidemiological unit* is affected by the entire sampling process, including sample selection, collection, handling and processing, as well as the actual laboratory test performance.

7. **Multiple sources of information**

Where multiple different data sources providing evidence of freedom from *infection* exist, each of these data sources may be analysed accordingly. The resulting estimates of the confidence in each data source may be combined to provide an overall level of confidence for the combined data sources.

The methodology used to combine the estimates from multiple data sources:

- *a)* should be scientifically valid, and fully documented, including references to published material; and
- *b)* should, where possible, take into account any lack of statistical independence between different data sources.

*Surveillance* information gathered from the same country, zone or compartment at different times (e.g. repeated annual surveys) may provide cumulative evidence of animal health status. Such evidence gathered over time may
be combined to provide an overall level of confidence. However, a single larger survey, or the combination of data collected during the same time period from multiple random or non-random sources, may be able to achieve the same level of confidence in a shorter period of time.

Analysis of surveillance information gathered intermittently or continuously over time should, where possible, incorporate the time of collection of the information to take into account the decreased value of older information. The sensitivity, specificity and completeness of data from each source should also be taken into account for the final overall confidence level estimation.

8. Sampling

The objective of sampling from a population is to select a subset of units from the population that is representative of the population with respect to the characteristic of interest (in this case, the presence or absence of infection). The survey design may involve sampling at several levels. For sampling at the level of the epidemiological units or higher units, a formal probability sampling (e.g. simple random sampling) method should be used. Sampling should be carried out in such a way as to provide the best likelihood that the sample will be representative of the population, within the practical constraints imposed by different environments and production systems.

When sampling below the level of the epidemiological unit (e.g. individual animal), the sampling method used should provide the best practical chance of generating a sample that is representative of the population of the chosen epidemiological unit. Collecting a truly representative sample of individual animals (whether from a pond, cage or fishery) is often very difficult. To maximise the chance of finding infection, the aim should be to bias the sampling towards infected animals, e.g. selecting moribund animals, life stages with a greater chance of active infection, etc.

Biased sampling in this context involves sampling from a defined study population that has a different probability of infection than the target population of which it is a subpopulation. Once the study population has been identified, the objective is still to select a representative sample from this subpopulation.

The sampling method used at all levels should be fully documented and justified.

9. Sample size

The number of units to be sampled from a population should be calculated using a statistically valid technique that takes at least the following factors into account:

- the sensitivity and specificity of the diagnostic test, or test system;
- the design prevalence (or prevalences where a multi-stage design is used);
- the level of confidence that is desired of the survey results.

Additionally, other factors may be considered in sample size calculations, including (but not limited to):

- the size of the population (but it is acceptable to assume that the population is infinitely large);
- the desired power of the survey;
- uncertainty about sensitivity and specificity.

The specific sampling requirements will need to be tailor-made for each individual disease, taking into account its characteristics and the specificity and sensitivity of the accepted testing methods for detecting the pathogenic agent in host populations.

FreeCalc is a suitable software for the calculation of sample sizes at varying parameter values. The table below provides examples of sample sizes generated by the software for a type I and type II error of 5% (i.e. 95% confidence and 95% statistical power). However, this does not mean that a type 1 and type 2 error of 0.05 should always be used. For example, using a test with sensitivity and specificity of 99%, 528 units should be sampled. If nine or less of those units test positive, the population can still be considered free of the disease at a design prevalence of 2% provided that all efforts are made to ensure that all presumed false positives are indeed false. This means that there is a 95% confidence that the prevalence is 2% or lower.

In the case in which the values of Se and Sp are not known (e.g. no information is available in the disease-specific chapter in the Aquatic Manual), they should not automatically be assumed to be 100%. All positive results should be included and discussed in any report regarding that particular survey and all efforts should be made to ensure that all presumed false positives are indeed false.
10. **Quality assurance**

Surveys should include a documented quality assurance system, to ensure that field and other procedures conform to the specified survey design. Acceptable systems may be quite simple, as long as they provide verifiable documentation of procedures and basic checks to detect significant deviations of procedures from those documented in the survey design.

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<th>Specificity (%)</th>
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Specific requirements for complex non-survey data sources for freedom from disease

Data sources that provide evidence of freedom from infection, but are not based on structured population-based surveys may also be used to demonstrate freedom, either alone or in combination with other data sources. Different methodologies may be used for the analysis of such data sources, but the methodology should comply with the provisions of this chapter. The approach used should, where possible, also take into account any lack of statistical independence between observations.

Analytical methodologies based on the use of step-wise probability estimates to describe the surveillance system may determine the probability of each step either by:

1) the analysis of available data, using a scientifically valid methodology; or where no data are available,
2) the use of estimates based on expert opinion, gathered and combined using a formal, documented and scientifically valid methodology.

Where there is significant uncertainty and/or variability in estimates used in the analysis, stochastic modelling or other equivalent techniques should be used to assess the impact of this uncertainty and/or variability on the final estimate of confidence.

Surveillance for distribution and occurrence of disease

Surveillance to determine distribution and occurrence of disease or of other relevant health related events is widely used to assess the prevalence and incidence of selected disease as an aid to decision-making, for example implementation of control and eradication programmes. It also has relevance for the international movement of animals and products when movement occurs among infected countries.

In contrast to surveillance to demonstrate freedom from disease, surveillance for the distribution and occurrence of disease is usually designed to collect data about a number of variables of animal health relevance, for example:

- prevalence or incidence of disease in wild or cultured animals;
- morbidity and mortality rates;
- frequency of disease risk factors and their quantification;
- frequency distribution of variables in epidemiological units;
- frequency distribution of the number of days elapsing between suspicion of disease and laboratory confirmation of the diagnostic and/or to the adoption of control measures;
- farm production records, etc.

This article describes surveillance to estimate parameters of disease occurrence.
1. Objectives

The objective of this kind of surveillance system is to contribute on an on-going basis evidence to assess the occurrence and distribution of disease or infection in a particular country, zone or compartment. This will provide information for domestic disease control programmes and relevant disease occurrence information to be used by trading partners for qualitative and quantitative risk assessment.

A single such survey can contribute evidence adding to an on-going collection of health data.

2. Population

The population of epidemiological units should be clearly defined. The target population consists of all individuals of all species susceptible to the disease in a country, zone or compartment to which the surveillance results apply. Some local areas within a region may be known to be free of the disease of concern, allowing resources to be concentrated on known positive areas for greater precision of prevalence estimates and only verification of expected 0 prevalence areas.

The design of the survey will depend on the size and structure of the population being studied. If the population is relatively small and can be considered to be homogenous with regards to risk of infection, a single-stage survey can be used.

In larger populations where a sampling frame is not available, or when there is a likelihood of clustering of disease, multi-stage sampling is required. For example, a multi-stage sampling process may involve sampling of farms or villages followed by sampling of fish from selected ponds within the sampled farms/villages.

In the case of a complex (e.g. multi-level) population structure, multi-level sampling may be used and the data analysed accordingly.

3. Sources of evidence

Surveillance data may originate from a number of different sources, including:

a) population-based surveys using one or more tests to detect the agent;

b) other non-random sources of data, such as:
   i) sentinel sites;
   ii) disease notifications and laboratory investigation records;
   iii) academic and other scientific studies;

c) a knowledge of the biology of the agent, including environmental, host population distribution, known geographical distribution, vector distribution and climatic information;

d) history of imports of potentially infected material;

e) biosecurity measures in place;

f) any other sources of information that provide contributory evidence regarding disease or infection in the country, zone or compartment.

The sources of evidence should be fully described. In the case of a structured survey, this should include a description of the sampling strategy used for the selection of units for testing. For complex surveillance systems, a full description of the system is required including consideration of any biases that may be inherent in the system. Evidence to support changes in prevalence/incidence of endemic disease should be based on valid, reliable methods to generate precise estimates with known error.

4. Statistical methodology

Analysis of survey data should be in accordance with the provisions of this chapter and should consider the following factors:

a) the survey design;

b) the sensitivity and specificity of the test, or test system;

c) the results of the survey.

For surveillance systems used to describe disease patterns, the purpose is to estimate prevalence or incidence with confidence intervals or probability intervals. The magnitude of these intervals expresses the precision of the estimates and is related to sample size. Narrow intervals are desirable but will require larger sample sizes and more dedication of resources. The precision of the estimates and the power to detect differences in prevalence between populations or between time points depends not only on sample size, but also on the actual value of the prevalence in the population or the actual difference. For this reason, when designing the surveillance system, a prior estimate/assumption of expected prevalence or expected difference in prevalence should be made.

For the purpose of describing disease occurrence, measures of animal unit, time and place can be calculated for an entire population and specific time period, or for subsets defined by host characteristics (e.g. age-specific
incidence). Incidence estimation requires on-going surveillance to detect new cases in a specified time period while prevalence is the estimated proportion of infected individuals in a population at a given time point. The estimation process should consider test sensitivity and specificity.

Statistical analysis of surveillance data often requires assumptions about population parameters or test characteristics. These are usually based on expert opinion, previous studies on the same or different populations, expected biology of the agent, information contained in the disease-specific chapter of the Aquatic Manual, and so on. The uncertainty around these assumptions should be quantified and considered in the analysis (e.g. in the form of prior probability distributions in a Bayesian setting).

When surveillance objectives are to estimate prevalence/incidence or changes in disease patterns, statistical analysis should account for sampling error. Analytic methods should be thoroughly considered and consultation with biostatistician/quantitative epidemiologist consulted beginning in the planning stages and continued throughout the programme.

5. Clustering of infection

Infection in a country, zone or compartment usually clusters rather than being uniformly distributed through a population. Clustering may occur at a number of different levels (e.g. a cluster of moribund fish in a pond, a cluster of ponds in a farm, or a cluster of farms in a zone). Except when dealing with demonstrably homogenous populations, surveillance should take this clustering into account in the design and the statistical analysis of the data, at least at what is judged to be the most significant level of clustering for the particular animal population and infection. For endemic diseases, it is important to identify characteristics of the population which contribute to clustering and thus provide efficiency in disease investigation and control.

6. Test characteristics

All surveillance involves performing one or more tests for evidence of the presence of current or past infection, ranging from detailed laboratory examinations to farmer observations. The performance level of a test at the population level is described in terms of its sensitivity and specificity. Imperfect sensitivity and/or specificity impact on the interpretation of surveillance results and should be taken into account in the analysis of surveillance data. For example, in populations with low prevalence of infection, a large proportion of positive tests may be false unless the tests used have perfect specificity. To ensure detection in such instances, a highly sensitive test is frequently used for initial screening and then confirmed with highly specific tests.

All calculations should take the performance level (sensitivity and specificity) of any tests used into account. The values of sensitivity and specificity used for calculations should be specified, and the method used to determine or estimate these values should be documented. Test sensitivity and specificity can be different when applied to different populations and testing scenarios. For example, test sensitivity may be lower when testing carrier animals with low level infections compared to moribund animals with clinical disease. Alternatively, specificity depends on the presence of cross-reacting agents, the distribution of which may be different under different conditions or regions. Ideally, test performance should be assessed under the conditions of use otherwise increased uncertainty exists regarding their performance. In the absence of local assessment of tests, values for sensitivity and/or specificity for a particular test that are specified in the Aquatic Manual may be used but the increased uncertainty associated with these estimates should be incorporated into the analysis of results.

Pooled testing involves the pooling of specimens from multiple individuals and performing a single test on the pool. Pooled testing is an acceptable approach in many situations. Where pooled testing is used, the results of testing should be interpreted using sensitivity and specificity values that have been determined or estimated for that particular pooled testing procedure and for the applicable pool sizes being used. Analysis of the results of pooled testing should, where possible, be performed using accepted, statistically based methodologies, which should be fully documented, including published references.

Test results from surveillance for endemic disease will provide estimates of apparent prevalence (AP). Using diagnostic sensitivity (DSe) and diagnostic specificity (DSp), true prevalence (TP) should be calculated with the following formula:

\[
TP = \frac{(AP + DSp - 1)}{(DSe + DSp - 1)}
\]

In addition, it should be remembered that different laboratories may obtain conflicting results for various tests, host, or procedure-related reasons. Therefore, sensitivity and specificity parameters should be validated for the particular laboratory and process.
7. Multiple sources of information

Where multiple different data sources providing information on infection or disease are generated, each of these data sources may be analysed and presented separately.

Surveillance information gathered from the same country, zone or compartment at different times and similar methodology (e.g. repeated annual surveys) may provide cumulative evidence of animal health status and changes. Such evidence gathered over time may be combined (e.g. using Bayesian methodology) to provide more precise estimates and details of disease distribution within a population.

Apparent changes in disease occurrence of endemic diseases may be real or due to other factors influencing detection proficiency.

8. Sampling

The objective of sampling from a population is to select a subset of units from the population that is representative of the population with respect to the characteristic of interest (in this case, the presence or absence of infection). The survey design may involve sampling at several levels. For sampling at the level of the epidemiological units or higher units, a formal probability sampling (e.g. simple random sampling) method should be used. Sampling should be carried out in such a way as to provide the best likelihood that the sample will be representative of the population, within the practical constraints imposed by different environments and production systems.

When sampling below the level of the epidemiological unit (e.g. individual animal), the method used should be probability-based sampling. Collecting a true probability-based sample is often very difficult and care should therefore be taken in the analysis and interpretation of results obtained using any other method, the danger being that inferences could not be made about the sampled population.

The sampling method used at all levels should be fully documented and justified.

9. Sample size

The number of units to be sampled from a population should be calculated using a statistically valid technique that takes at least the following factors into account:

- the sensitivity and specificity of the diagnostic test (single or in combination);
- expected prevalence or incidence in the population (or prevalences/incidences where a multi-stage design is used);
- the level of confidence that is desired of the survey results;
- the precision desired (i.e. the width of the confidence or probability intervals).

Additionally, other factors may be considered in sample size calculations, including (but not limited to):

- the size of the population (but it is acceptable to assume that the population is infinitely large);
- uncertainty about sensitivity and specificity.

The specific sampling requirements will need to be tailor-made for each individual disease, taking into account its characteristics and the specificity and sensitivity of the accepted testing methods for detecting the pathogenic agent in host populations.

A number of software packages, e.g. Survey Tool Box (www.aciar.gov.au; www.ausvet.com.au), WinPEPI (www.sagebrushpress.com/pepibook.html) can be used for the calculation of sample sizes.

In the case in which the values of Se and Sp are not known (e.g. no information is available in the disease-specific chapter in the Aquatic Manual), they should not automatically be assumed to be 100%. Assumed values should be produced in consultation with subject-matter experts.

10. Quality assurance

Surveys should include a documented quality assurance system, to ensure that field and other procedures conform to the specified survey design. Acceptable systems may be quite simple, as long as they provide verifiable documentation of procedures and basic checks to detect significant deviations of procedures from those documented in the survey design.
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Article 1.4.11.

Examples of surveillance programmes

The following examples describe surveillance systems and approaches to the analysis of evidence for demonstrating freedom from disease. The purpose of these examples is:

– to illustrate the range of approaches that may be acceptable;
– to provide practical guidance and models that may be used for the design of specific surveillance systems; and
– to provide references to available resources that are useful in the development and analysis of surveillance systems.

While these examples demonstrate ways in which freedom from disease may be successfully demonstrated, they are not intended to be prescriptive. Countries are free to use different approaches, as long as they meet the requirements of this chapter.

The examples deal with the use of surveys and are designed to illustrate different survey designs, sampling schemes, the calculation of sample size, and analysis of results. It is important to note that alternative approaches to demonstrating freedom using complex non-survey-based data sources are also currently being developed and may soon be published.

1. Example 1. – One-stage structured survey (farm certification)

a) Context

A freshwater aquaculture industry raising fish in tanks has established a farm certification scheme. This involves demonstrating farm-level freedom from a particular (hypothetical) disease (Disease X). The disease does not spread very quickly, and is most common during the winter months, with adult fish at the end of the production cycle being most severely affected. Farms consist of a number of grow-out tanks, ranging from 2 to 20, and each tank holds between 1,000 and 5,000 fish.

b) Objective

The objective is to implement surveillance that is capable of providing evidence that an individual farm is free from Disease X. (The issue of national or zone freedom, as opposed to farm freedom, is considered in the next example.)

c) Approach

The accreditation scheme establishes a set of standard operating procedures and requirements for declaration of freedom, based on the recommendations given in this chapter. These require farms to undertake a survey capable of producing 95% confidence that the disease would be detected if it were present. Once farms have been surveyed without detecting disease, they are recognised as free, as long as they maintain a set of minimum biosecurity standards. These standards are designed to prevent the introduction of Disease X into the farm (through the implementation of controls specific to the method of spread of that disease) and to ensure that the disease would be detected rapidly if it were to enter the farm (based on evidence of adequate health record keeping and the prompt investigation of unusual disease events). The effective implementation of these biosecurity measures is evaluated with annual on-farm audits conducted by independent auditors.

d) Survey standards

Based on the recommendations given in this chapter, a set of standards are established for the conduct of surveys to demonstrate freedom from infection with causative agent of Disease X. These standards include:

i) The level of confidence required of the survey is 95% (i.e. Type I error = 5%).

ii) The power of the survey is arbitrarily set at 95% (i.e. Type II error = 5%, which means that there is a 5% chance of concluding that a non-diseased farm is infected).

iii) The target population is all the fish on the farm. Due to the patterns of disease in this production system, in which only fish in the final stages of grow-out, and only in winter are affected, the study population is defined as grow-out fish during the winter months.

iv) The issue of clustering is considered. As fish are grouped into tanks, this is the logical level at which to consider clustering. However, when a farm is infected, the disease often occurs in multiple tanks, so there is little evidence of strong clustering. Also, the small number of tanks on a single farm means that
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it is difficult to define a design prevalence at the tank level (i.e. the proportion of infected tanks that the survey should be able to detect on the farm). For these reasons, it is decided to treat the entire grow-out population of each farm as a single homogenous population.

v) Stratification is also considered. In order to ensure full representation, it is decided to stratify the sample size by tank, proportional to the population of each tank.

vi) The design prevalence at the animal level is determined based on the epidemiology of the disease. The disease does not spread quickly; however, in the defined target population, it has been reported to affect at least 10% of fish if the population is infected. In order to take the most conservative approach, an arbitrarily low design prevalence of 2% is used. A prevalence of 10% may have been used (and would result in a much smaller sample size), but the authorities were not convinced by the thought that the population could still be infected at a level of say 5%, and disease still not be detected.

vii) The test used involves destructive sampling of the fish, and is based on an antigen-detection enzyme-linked immunosorbent assay (ELISA). Disease X is present in some parts of the country (hence the need for a farm-level accreditation programme). This has provided the opportunity for the sensitivity and the specificity of the ELISA to be evaluated in similar populations to those on farms. A recent study (using a combination of histology and culture as a gold standard) estimated the sensitivity of the ELISA to be 98% (95% confidence interval 96.7–99.2%), and the specificity to be 99.4% (99.2–99.6%). Due to the relatively narrow confidence intervals, it was decided to use the point estimates of the sensitivity and specificity rather than complicate calculations by taking the uncertainty in those estimates into account.

e) Sample size

The sample size required to meet the objectives of the survey is calculated to take the population size, the test performance, the confidence required and the design prevalence into account. As the population of each farm is relatively large, differences in the total population of each farm have little effect on the calculated sample size. The other parameters for sample size calculation are fixed across all farms. Therefore, a standard sample size (based on the use of this particular ELISA, in this population) is calculated. The sample size calculations are performed using the FreeCalc software. Based on the parameters listed above, the sample size required is calculated to be 410 fish per farm. In addition, the programme calculates that, given the imperfect specificity, it is still possible for the test to produce up to five false-positive reactors from an uninfected population using this sample size. The authorities are not comfortable with dealing with false-positive reactors, so it is decided to change the test system to include a confirmatory test for any positive reactors. Culture is selected as the most appropriate test, as it has a specificity that is considered to be 100%. However, its sensitivity is only 90% due to the difficulty of growing the organism.

As two tests are now being used, the performance of the test system should be calculated, and the sample size recalculated based on the test system performance.

Using this combination of tests (in which a sample is considered positive only if it tests positive to both tests), the specificity of the combined two tests can be calculated by the formula:

\[ Sp_{combined} = Sp_1 \times Sp_2 - (Sp_1 \times Sp_2) \]

which produces a combined specificity of \(1 + 0.994 - (1 \times 0.994) = 100\%\).

The sensitivity may be calculated by the formula:

\[ Spe_{combined} = Se_1 \times Se \]

which produces a combined sensitivity of \(0.9 \times 0.98 = 88.2\%\).

These new values are used to calculate the survey sample size yielding a result of 169 fish. It is worth noting that attempts to improve the performance of a test (in this case increase specificity) generally result in a decrease in the performance of the other aspect of the test performance (sensitivity in this example).
However, in this case, the loss of sensitivity is more than compensated for by the decreased sample size due to the improved specificity.

It is also worth noting that, when using a test system with 100% specificity, the effective power of the survey will always be 100%, regardless of the figure used in the design. This is because it is not possible to make a Type II error, and conclude that the farm is infected when it is not.

A check of the impact of population size on the calculated sample size is worthwhile. The calculated sample size is based on an infinitely large population. If the population size is smaller, the impact on sample size is shown in the following table:

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<tr>
<td>2,000</td>
<td>163</td>
</tr>
<tr>
<td>5,000</td>
<td>166</td>
</tr>
<tr>
<td>10,000</td>
<td>169</td>
</tr>
</tbody>
</table>

Based on these calculations, it is clear that, for the population sizes under consideration, there is little effect on the sample size. For the sake of simplicity, a standard sample size of 169 is used, regardless of the number of grow-out fish on the farm.

f) Sampling

The selection of individual fish to include in the sample should be done in such a manner as to give the best chance of the sample being representative of the study population. A fuller description of how this may be achieved under different circumstances is provided in Survey Toolbox3. An example of a single farm will be used to illustrate some of the issues.

One farm has a total of eight tanks, four of which are used for grow-out. At the time of the survey (during winter), the four grow-out tanks have 1,850, 4,250, 4,270 and 4,880 fish, respectively, giving a total population of 15,250 grow-out fish.

Simple random sampling from this entire population is likely to produce sample sizes from each tank roughly in proportion to the number of fish in each tank. However, proportional stratified sampling will guarantee that each tank is represented in proportion. This simply involves dividing the sample size between tanks in proportion to their population. The first tank has 1,850 fish out of a total of 15,250, representing 12.13%. Therefore, 12.13% of the sample (21 fish) should be taken from the first tank. Using a similar approach the sample size for the other three tanks is 47, 47 and 54 fish, respectively.

Once the sample for each tank is determined, the problem remains as to how to select 21 fish from a tank of 1,850 so that they are representative of the population. Several options exist.

1) If the fish can be handled individually, random systematic sampling may be used. For example, samples can be collected at harvest or during routine management activities involving handling the fish (such as grading or vaccination).

   If fish are handled, systematic sampling simply involves selecting a fish at regular intervals. For instance, to select 21 from 1,850, the sampling interval should be 1,850/21 = 88. This means that every 88th fish from the tank should be sampled. To ensure randomness, it is good practice to use a random number between 1 and 88 (in this case) to select the first fish (e.g., using a random number table), and then select every 88th fish after that.

2) If fish cannot be handled individually (by far the most common, and more difficult, circumstance) then the fish to be sampled should be captured from the tanks. Fish should be captured in the most efficient and practical way possible; however, every effort should be made to try to ensure that the sample is representative. In this example, a dip net is the normal method used for capturing fish. Using a dip net, convenience sampling would involve capturing 21 fish by repeatedly dipping at one spot and capturing the easiest fish (perhaps the smaller ones). This approach is strongly discouraged. One method of increasing the representativeness is to sample at different locations in the tank — some at one end, some at either side, some at the other end, some in the middle, some close to the edge. Additionally, if there are differences among the fish, an attempt should be made to capture fish in such a way as to give different groups of fish a chance of being caught (i.e., do not just try to catch the small ones, but include big ones as well).

   This method of collecting a sample is far from the ideal of random sampling, but due to the practical difficulties of implementing random sampling of individual fish, this approach is acceptable, as long as...
the efforts made to increase the representativeness of the sample are both genuine and fully documented.

g) Testing

Specimens are collected, processed and tested in accordance with standardised procedures developed under the certification programme and designed to meet the requirements of the Aquatic Manual. The testing protocol dictates that any specimens that test positive to ELISA be submitted for culture, and that any positive culture results indicate a true positive specimen (i.e. that the farm is not free from disease). It is important that this protocol be adhered to exactly. If a positive culture is found, then it is not acceptable to retest it, unless further testing is specified in the original testing protocol, and the impact of such testing accounted for in the test system sensitivity and specificity estimates (and therefore the sample size).

h) Analysis

If the calculated sample size of 169 is used, and no positive reactors are found, then the survey will have a confidence of 95%. This can be confirmed by analysing the results using the FreeCalc software mentioned above (which reports a confidence level of 95.06%).

It may happen in some cases that the survey is not conducted exactly as planned, and the actual sample size is less than the target sample size. However, the size of the farm may also be smaller. In these cases, it is advisable to analyse the farm data on a farm-by-farm basis. For example, if only 165 specimens were collected from a farm with only 2,520 fish, the resulting confidence would still be 95%. If only 160 fish were collected, the confidence is only 94.5%. If a rigid target of 95% confidence is used, then this survey would fail to meet that target and more evidence would be required.

2. Example 2 – Two-stages structured survey (national freedom)

a) Context

A country aims to declare freedom from Disease Y of crustaceans. The industry in this country is based largely on small-holder ponds, grouped closely together in and around villages. The disease is reasonably highly contagious, and causes mass mortality mid to late in the production cycle, with affected animals becoming moribund and dying in a matter of days. Affected animals show few characteristic signs, but an infected pond will almost invariably break down with mass mortality unless harvested beforehand. It is more common in late summer, but can occur at any time of year. It also occurs occasionally early in the production cycle. In this country, there are some limitations to the availability of laboratory facilities and the transport infrastructure. However, there is a relatively large government structure, and a comprehensive network of fisheries officers.

b) Objective

The objective is to establish national freedom from Disease Y. The surveillance system should meet the requirements of this chapter, but should also be able to be practically implemented in this small-holder production system.

c) Approach

The aquaculture authorities decide to use a survey to gather evidence of freedom, using a two-stage survey design (sampling villages at the first level, and ponds at the second). Laboratory testing of specimens from a large number of farms is not considered feasible, so a combined test system is developed to minimise the need for expensive laboratory tests.

The unit of observation and analysis is, in this case, the pond, rather than the individual animal. This means that the diagnostic is being made at the pond level (an infected pond or a non-infected pond) rather than at the animal level.

The survey is therefore a survey to demonstrate that no villages are infected (using a random sample of villages and making a village-level diagnostic). The test used to make a village-level diagnostic is, in fact, another survey, this time to demonstrate that no ponds in the village are affected. A test is then performed at the pond level (farmer observation followed, if necessary, by further laboratory testing).
d) Survey standards

i) The confidence to be achieved by the survey is 95%. The power is set at 95% (but is likely to be virtually 100% if the test system used achieves nearly 100% specificity, as demonstrated in the previous example).

ii) The target population is all ponds stocked with shrimp in the country during the study period. The study population is the same, except that those remote areas to which access is not possible are excluded. As outbreaks can occur at any time of year, and at any stage of the production cycle, it is decided not to further refine the definition of the population to target a particular time or age.

iii) Three tests are used. The first is farmer observation, to determine if mass mortality is occurring in a particular pond. If a pond is positive to the first test (i.e. mass mortality is detected), a second test is applied. The second test used is polymerase chain reaction (PCR). Cases positive to PCR are further tested using transmission experiments.

iv) Farmer observation can be treated as a test just like any other. In this case, the observation of mass mortality is being used as a test for the presence of Disease Y. As there are a variety of other diseases that are capable of causing mass mortality, the test is not very specific. On the other hand, it is quite unusual for Disease Y to be present, and not result in mass mortality, so the test is quite sensitive. A standard case definition is established for mass mortality (for instance, greater than 20% of the pond’s population of shrimp observed dead in the space of less than one week). Based on this definition, farmers are able to ‘diagnose’ each pond as having mass mortality. Some farmers may be over-sensitive and decide that mass mortality is occurring when only a small proportion of shrimp are found dead (false positives, leading to a decrease in specificity) while a small number of others fail to recognise the mortalities, decreasing sensitivity.

In order to quantify the sensitivity and specificity of farmer observation of mass mortalities, as a test for Disease Y, a separate study is carried out. This involves both a retrospective study of the number of mass mortality events in a population that is thought to be free from disease, as well as a study of farmers presented with a series of mortality scenarios, to assess their ability to accurately identify a pond with mass mortality. By combining these results, it is estimated that the sensitivity of farmer-reported mass mortalities as a test for Disease Y is 87% while the specificity is 68%.

v) When a farmer detects a pond with mass mortality, specimens are collected from moribund shrimp following a prescribed protocol. Tissue samples from 20 shrimp are collected, and pooled for PCR testing. In the laboratory, the ability of pooled PCR to identify a single infected animal in a pool of 20 has been studied, and the sensitivity of the procedure is 98.6%. A similar study of negative specimens has shown that positive results have occasionally occurred, probably due to laboratory contamination, but maybe also because of the presence of non-viable genetic material from another source (shrimp-based feed stuffs are suspected). The specificity is therefore estimated at 99%.

vi) Published studies in other countries have shown that the sensitivity of transmission tests, the third type of test to be used, is 95%, partly due to variability in the load of the agent in inoculated material. The specificity is agreed to be 100%.

vii) Based on these figures, the combined test system sensitivity and specificity are calculated using the formulae presented in Example 1, first with the first two tests, and then with the combined effect of the first two tests and the third test. The result is a sensitivity of 81.5% and a specificity of 100%.

viii) The design prevalence should be calculated at two levels. First, the pond-level design prevalence (the proportion of ponds in a village that would be infected if disease were present) is determined. In neighbouring infected countries, experience has shown that ponds in close contact with each other are quickly infected. It is unusual to observe an infected village with fewer than 20% of ponds infected. Conservatively, a design prevalence of 5% is used. The second value for design prevalence applies at the village level, or the proportion of infected villages that could be identified by the survey. As it is conceivable that the infection may persist in a local area without rapid spread to other parts of the country, a value of 1% is used. This is considered to be the lowest design prevalence value for which a survey can be practically designed.
ix) The population of villages in the country is 65,302, in accordance with official government records. Those with shrimp ponds number 12,890, based on records maintained by the aquaculture authorities. These are generated through a five-yearly agricultural census, and updated annually based on reports of fisheries officers. There are no records available of the number of ponds in each of these villages.

e) Sample size

Sample size is calculated for the two levels of sampling, first the number of villages to be sampled and then the number of ponds to be sampled. The number of villages to be sampled depends on the sensitivity and the specificity of the test used to classify villages as infected or not infected. As the ‘test’ used in each village is really just another survey, the sensitivity is equal to the confidence and the specificity is equal to the power of the village-level survey. It is possible to adjust both confidence and power by changing the sample size in the village survey (number of ponds examined), which means that we can determine, within certain limits, what sensitivity and specificity we achieve.

This allows a flexible approach to sample size calculation. If a smaller first-stage sample size is desired (a small number of villages), a high sensitivity and specificity are needed, which means that the number of ponds in each village that need to be examined is larger. A smaller number of ponds will result in lower sensitivity and specificity, requiring a larger number of villages. The approach to determining the optimal (least cost) combination of first- and second-stage sample sizes is described in Survey Toolbox.

A further complication is presented by the fact that each village has a different number of ponds. In order to achieve the same (or similar) confidence and power (sensitivity and specificity) for each village, a different sample size may be required. The authorities choose to produce a table of sample sizes for the number of ponds to sample in each village, based on the total ponds in each village.

An example of one possible approach to determining the sample size follows:

The target sensitivity (confidence) achieved by each village-level survey is 95%. The target specificity is 100%. Using the FreeCalc software, with a design prevalence of 1% (the survey is able to detect disease if 1% or more villages are infected), the first-stage sample size is calculated as 314 villages. Within each village, the test used is the combined test system described above with a sensitivity of 81.5% and a specificity of 100%. Based on these figures the following table is developed, listing the number of ponds that need to be sampled in order to achieve 95% sensitivity.

f) Sampling

First-stage sampling (selection of villages) is done using random numbers and a sampling frame based on the fisheries authorities list of villages with shrimp ponds. The villages are listed on a spreadsheet with each village numbered from 1 to 12,890. A random number table (such as that included in Survey Toolbox) or software designed for the generation of random numbers (such as EpiCalc 2) is used.

The second stage of sampling involves random selection of ponds within each village. This requires a sampling frame, or list of each pond in the village. The fisheries authorities use trained local fisheries officers to coordinate the survey. For each selected village, the officer visits the village and convenes a meeting of all shrimp farmers. At the meeting, they are asked how many ponds they have and a list of farmers’ names and the number of ponds is compiled. A simple random sample of the appropriate number of ponds (between 29 and 70, from the table above, depending on the number of ponds in the village) is selected from this list. This is done either using software (such as Survey Toolbox’s Random Animal Programme), or manually with a random number table or decimal dice for random number selection. Details of this process are described in Survey Toolbox. This selection process identifies a particular pond in terms of the name of the owner, and the
sequence number amongst the ponds owned (e.g. Mr Smith’s 3rd pond). Identification of the actual pond is based on the owners own numbering system for the ponds.

<table>
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<tr>
<th>Population</th>
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<td>1,000</td>
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</table>

**g) Testing**

Once ponds have been identified, the actual survey consists of ‘testing those ponds’. In practice, this involves the farmers observing the ponds during one complete production cycle. The local fisheries officer makes weekly visits to each farmer to check if any of the selected ponds have suffered mass mortality. If any are observed (i.e. the first test is positive), 20 moribund shrimp are collected for laboratory examination (first PCR, and then, if positive, transmission experiments).

**h) Analysis**

Analysis is performed in two stages. First, the results from each village are analysed to ensure that they meet the required level of confidence. If the target sample size is achieved (and only negative results obtained), the confidence should be 95% or greater in each village. At the second stage, the results from each village are analysed to provide a country level of confidence. Again, if the target sample size (number of villages) is achieved, this should exceed 95%.
3. Example 3. – Spatial sampling and the use of tests with imperfect specificity

a) Context

A country has an oyster culture industry, based primarily on rack culture of oysters in 23 estuaries distributed along the coastline. In similar regions in other countries, Disease Z causes mortalities in late summer/early autumn. During an outbreak a high proportion of oysters are affected; however, it is suspected that the agent may be present at relatively low prevalence in the absence of disease outbreaks.

b) Objective

The national authorities wish to demonstrate national freedom from Disease Z. If the disease should be detected, a secondary objective of the survey is to collect adequate evidence to support zoning at the estuary level.

c) Approach

The authorities conclude that clinical surveillance for disease outbreaks is inadequate because of the possibility of low level subclinical infections. It is therefore decided to base surveillance on a two-stage survey, in which sampled oysters are subjected to laboratory testing. The first stage of the survey is the selection of estuaries. However, due to the objective of providing evidence for zoning (should disease be found in any of the estuaries), it is decided to use a census approach and sample every estuary. In essence this means that there will be 23 separate surveys, one for each estuary. A range of options for sampling oysters are considered, including sampling at harvest or marketing, or using farms (oyster leases) as a level of sampling or stratification. However, the peak time of activity of the agent does not correspond to the harvest period, and the use of farms would exclude the significant numbers of wild oysters present in the estuaries. It is therefore decided to attempt to simulate simple random sampling from the entire oyster population in the estuary, using a spatial sampling approach.

d) Survey standards

i) The target population is all of the oysters in each of the estuaries. The study population is the oysters present during the peak disease-risk period in late summer early autumn. Wild and cultured oysters are both susceptible to disease, and may have associated with them different (but unknown) risks of infection. They are therefore both included in the study population. As will be described below, sampling is based on mapping. Therefore the study population can more accurately be described as that population falling within those mapped areas identified as oyster habitats.

ii) A design prevalence value is only required at the oyster level (as a census is being used at the estuary level). While the disease is often recognised with a very high prevalence during outbreaks, a low value is used to account for the possibility of persistence of the agent in the absence of clinical signs. A value of 2% is selected.

iii) The test used is histopathology with immuno-staining techniques. This test is known to produce occasional false-positive results due to nonspecific staining, but is very sensitive. Published studies indicate values of 99.1% for sensitivity and 98.2% for specificity. No other practical tests are available. This means that it is not possible to definitively differentiate false positives from true positives, and that in a survey of any size, a few false positives are expected (i.e. 1.8%).

iv) The confidence is set at 95% and the power at 80%. In the previous examples, due to the assumed 100% specificity achieved by use of multiple tests, the effective power was 100%. In this case, with imperfect specificity, there will be a risk of falsely concluding that a healthy estuary is infected, so the power is not 100%. The choice of a relatively low figure (80%) means that there is a one in five chance of falsely calling an estuary infected when it is not infected, but it also dramatically decreases the survey costs, through a lower sample size.
Chapter 1.4.- Aquatic animal health surveillance

**e) Sample size**

Based on the assumption that the sampling procedure will mimic simple random sampling, the sample size (number of oysters to sample per estuary) can be calculated with FreeCalc. The population size (number of oysters per estuary) is assumed to be very large. The calculated sample size, using the sensitivity, specificity and design prevalence figures given above, is 450. FreeCalc also reports that, based on this sample size and the specificity of the test, it is possible to get ten or fewer false-positive test results, and still conclude that the population is free from disease. This is because, if the population were infected at 2% or greater, the anticipated number of positive reactors from a sample of 450 would be greater than 10. In fact, we would expect 9 true positives (450 × 2% × 99.1%) and 8 false positives (450 × 98% × 1.8%) or a total of 17 positives if the population were infected at a prevalence of 2%.

This illustrates how probability theory and adequate sample size can help differentiate between true- and false-positive results when there is no alternative but to use a test with imperfect specificity.

**f) Sampling**

The aim is to collect a sample of 450 oysters that represent an entire estuary. Simple random sampling depends on creating a sampling frame listing every oyster (not possible) and systematic sampling depends on being able to (at least conceptually) line up all the oysters (again, not possible). The authorities decide to use spatial sampling to approximate simple random sampling. Spatial sampling involves selecting random points (defined by coordinates), and then selecting oysters near the selected points. In order to avoid selecting many points with no oysters nearby, the estuary is first mapped (the fisheries authorities already have digital maps defining oyster leases available). To these maps areas with significant concentrations of wild oysters are also added, based on local expertise. Pairs of random numbers are generated such that the defined point falls within the defined oyster areas. Other schemes are considered (including using a rope marked at regular intervals, laid out on a lease to define a transect, and collecting an oyster adjacent to each mark on the rope) but the random coordinate approach is adopted.

Survey then visit each point by boat (using a GPS Global Positioning System unit to pinpoint the location). A range of approaches is available for selecting which oyster to select from a densely populated area, but it should involve some effort at randomness. Survey staff opt for a simple approach: when the GPS receiver indicates that the site has been reached, a pebble is tossed in the air and the oyster closest to the point where it lands is selected. Where oysters are arranged vertically (e.g. wild oysters growing up a post), a systematic approach is used to determine the depth of the oyster to select. First, an oyster at the surface, next, an oyster halfway down, and thirdly, an oyster as deep as can be reached from the boat.

This approach runs the risk of bias towards lightly populated areas, so an estimate of the relative density of oysters at each sampling point is used to weight the results (see Survey Toolbox for more details).

**g) Testing**

Specimens are collected, processed, and analysed following a standardised procedure. The results are classified as definitively positive (showing strong staining in a highly characteristic pattern, possibly with associated signs of tissue damage), probably positive (on the balance of probabilities, but less characteristic staining), and negative.

**h) Analysis**

The interpretation of the results when using a test with imperfect specificity is based on the assumption that, in order to conclude that the population is free from infection, any positive result identified is really a false positive. With a sample size of 450, up to 10 false positives may be expected while still concluding that the population is free from disease. However, if there is reasonable evidence that there is even a single true positive, then the population cannot be considered free. This is the reason for the classification of positive results into definitive and probable positives. If there are any definitive positives at all, the population in that estuary should be considered infected. The probable positives are consistent with false positives, and therefore up to ten may be accepted. Using FreeCalc the actual confidence achieved based on the number of (presumed) false positives detected can be calculated. For instance, if 8 ‘probably positive’ results were detected from an estuary, the confidence level for the survey would be 98.76%. On the other hand, if 15 ‘probably positive’ results were detected, the confidence is only 61.9%, indicating that the estuary is likely to be infected.
Discussion

Normally, it may be safely assumed that a surveillance system aimed at demonstrating freedom from disease is 100% specific. This is because any suspected occurrence of disease is investigated until a definitive decision can be made. If the conclusion is that the case is truly a case of disease, then there is no issue of declaring freedom – the disease is known to be present. This example presents a different situation where, due to lack of suitable tests, it is not possible for the surveillance system to be 100% specific. This may represent an unusual situation in practice, but illustrates that methods exist for dealing with this sort of problem. In practice, a conclusion that a country (or estuary) is free from infection, in the face of a small (but statistically acceptable) number of positive results, will usually be backed up by further evidence (such as the absence of clinical disease).

NB: FIRST ADOPTED IN 2008; MOST RECENT UPDATE ADOPTED IN 2016.

1 FreeCalc – Cameron, AR. Software for the calculation of sample size and analysis of surveys to demonstrate freedom from disease. Available for free download from http://www.ausvet.com.au
4 http://www.myatt.demon.co.uk/epicalc.htm
CHAPTER 1.5.

CRITERIA FOR LISTING SPECIES AS SUSCEPTIBLE TO INFECTION WITH A SPECIFIC PATHOGEN

Article 1.5.1.

The purpose of this chapter is to provide criteria for determining which species are listed as susceptible in Article 1.5.2. of each disease-specific chapter in the Aquatic Code.

Article 1.5.2.

Scope

Susceptibility may include clinical or non-clinical infection but does not include species that may carry the pathogenic agent without replication.

The decision to list a species as susceptible should be based on a finding that the evidence is definite. However, possible susceptibility of a species is also important information and this should also be included in Section 2.2.1. entitled «Susceptible host species» of the relevant disease-specific chapter of the Aquatic Manual.

Article 1.5.3.

Approach

A three-stage approach is outlined in this chapter to assess susceptibility of a species to infection with a specified pathogenic agent and is based on:

1) criteria to determine whether the route of transmission is consistent with natural pathways for the infection (as described in Article 1.5.4.);

2) criteria to determine whether the pathogenic agent has been adequately identified (as described in Article 1.5.5.);

3) criteria to determine whether the evidence indicates that presence of the pathogenic agent constitutes an infection (as described in Article 1.5.6.).

Article 1.5.4.

Stage 1: criteria to determine whether the route of transmission is consistent with natural pathways for the infection

The evidence should be classified as transmission through:

1) natural occurrence; includes situations where infection has occurred without experimental intervention e.g. infection in wild or farmed populations; or

2) non-invasive experimental procedures; includes cohabitation with infected hosts, infection by immersion or ingestion; or

3) invasive experimental procedure; includes injection, exposure to unnaturally high loads of pathogen, or exposure to stressors (e.g. temperature) not encountered in the host's natural or culture environment.

Consideration needs to be given to whether experimental procedures (e.g. inoculation, infectivity load) mimic natural pathways for disease transmission. Consideration should also be given to environmental factors as these may affect host resistance or transmission of the pathogen.
Article 1.5.5.

Stage 2: criteria to determine whether the pathogenic agent has been adequately identified

The pathogenic agent should be identified and confirmed in accordance with the methods described in Section 7 (corroborative diagnostic criteria) of the relevant disease chapter in the Aquatic Manual, or other methods that have been demonstrated to be equivalent.

Article 1.5.6.

Stage 3: criteria to determine whether the evidence indicates that presence of the pathogenic agent constitutes an infection

A combination of the following criteria should be used to determine infection (see Article 1.5.7.):

A. the pathogenic agent is multiplying in the host, or developing stages of the pathogenic agent are present in or on the host;
B. viable pathogenic agent is isolated from the proposed susceptible species, or infectivity is demonstrated by way of transmission to naive individuals;
C. clinical or pathological changes are associated with the infection;
D. the specific location of the pathogen corresponds with the expected target tissues.

The type of evidence to demonstrate infection will depend on the pathogenic agent and potential host species under consideration.

Article 1.5.7.

Outcomes of the assessment

The decision to list a species as susceptible should be based on a finding of definite evidence. Evidence should be provided for the following:

1) transmission has been obtained naturally or by experimental procedures that mimic natural pathways for the infection in accordance with Article 1.5.4.;

AND

2) the identity of the pathogenic agent has been confirmed in accordance with Article 1.5.5.;

AND

3) there is evidence of infection with the pathogenic agent in the suspect host species in accordance with criteria A to D in Article 1.5.6. Evidence to support criterion A alone is sufficient to determine infection. In the absence of evidence to meet criterion A, satisfying at least two of criteria B, C or D would be required to determine infection.

Article 1.5.8.

Species for which there is incomplete evidence for susceptibility

The decision to list a species as susceptible in Article 1.5.2. of each disease-specific chapter should be based on a finding that the evidence is definite.

However, where there is insufficient evidence to demonstrate susceptibility through the approach described in Article 1.5.3. because transmission does not mimic natural pathways of infection, or the identity of the pathogenic agent has not been confirmed, or infection is only partially supported, information will be included in the relevant disease-specific chapter in the Aquatic Manual.
Chapter 1.5.- Criteria for listing species as susceptible to infection with a specific pathogen

If there is insufficient evidence to demonstrate susceptibility of a species, the Competent Authority should assess the risk of spread of the pathogen under consideration, in accordance with the recommendations in Chapter 2.1., prior to the implementation of import health measures.

NB: FIRST ADOPTED IN 2014; MOST RECENT UPDATE ADOPTED IN 2016.
SECTION 2.
RISK ANALYSIS

CHAPTER 2.1.
IMPORT RISK ANALYSIS

Article 2.1.1.

Introduction

The importation of aquatic animals and aquatic animal products involves a degree of disease risk to the importing country. This risk may be represented by one or several diseases or infections.

The principal aim of import risk analysis is to provide importing countries with an objective and defensible method of assessing the disease risks associated with the importation of aquatic animals, aquatic animal products, aquatic animal genetic material, feedstuffs, biological products and pathological material. The principles and methods are the same whether the commodities are derived from aquatic and/or terrestrial animal sources. The analysis should be transparent. This is necessary so that the exporting country is provided with clear reasons for the imposition of import conditions or refusal to import.

Transparency is also essential because data are often uncertain or incomplete and, without full documentation, the distinction between facts and the analyst’s value judgements may blur.

This chapter provides recommendations and principles for conducting transparent, objective and defensible risk analyses for international trade. However, it cannot provide details on the means by which a risk analysis is carried out as the purpose of the Aquatic Code is simply to outline the necessary basic steps. The components of risk analysis are hazard identification, risk assessment, risk management and risk communication (Figure 1).

Fig. 1. The four components of risk analysis

The risk assessment is the component of the analysis that estimates the risks associated with a hazard. Risk assessments may be qualitative or quantitative. For many diseases, particularly for those diseases listed in the Aquatic Code where there are well developed internationally agreed standards, there is broad agreement concerning the likely risks. In such cases it is more likely that a qualitative assessment is all that is required. Qualitative assessment does not require mathematical modelling skills to carry out and so is often the type of assessment used for routine decision-making. No single method of import risk assessment has proven applicable in all situations, and different methods may be appropriate in different circumstances.
Chapter 2.1.- Import risk analysis

The process of import risk analysis on aquatic animals and aquatic animal products usually needs to take into consideration the results of an evaluation of the Aquatic Animal Health Services, zoning and compartmentalisation, and surveillance systems that are in place for monitoring aquatic animal health in the exporting country. These are described in separate chapters in the Aquatic Code.

Article 2.1.2.

Hazard identification

Hazard identification involves identifying the pathogenic agents that could potentially produce adverse consequences associated with the importation of a commodity.

The hazards identified would be those appropriate to the species being imported, or from which the commodity is derived, and which may be present in the exporting country. It is then necessary to identify whether each hazard is already present in the importing country, and whether it is a listed disease or is subject to control or eradication in that country and to ensure that import measures are not more trade restrictive than those applied within the country.

Hazard identification is a categorisation step, identifying biological agents dichotomously as hazards or not hazards. The risk assessment should be concluded if hazard identification fails to identify hazards associated with the importation.

The evaluation of the Aquatic Animal Health Services, surveillance and control programmes, and zoning and compartmentalisation systems are important inputs for assessing the likelihood of hazards being present in the aquatic animal population of the exporting country.

An importing country may decide to permit the importation using the appropriate sanitary standards recommended in the Aquatic Code, thus eliminating the need for a risk assessment.

Article 2.1.3.

Principles of risk assessment

1) Risk assessment should be flexible in order to deal with the complexity of real-life situations. No single method is applicable in all cases. Risk assessment should be able to accommodate the variety of aquatic animal commodities, the multiple hazards that may be identified with an importation and the specificity of each disease, detection and surveillance systems, exposure scenarios and types and amounts of data and information.

2) Both qualitative risk assessment and quantitative risk assessment methods are valid.

3) The risk assessment should be based on the best available information that is in accord with current scientific thinking. The assessment should be well documented and supported with references to the scientific literature and other sources, including expert opinion.

4) Consistency in risk assessment methods should be encouraged and transparency is essential in order to ensure fairness and rationality, consistency in decision-making and ease of understanding by all the interested parties.

5) Risk assessments should document the uncertainties, the assumptions made, and the effect of these on the final risk estimate.

6) Risk increases with increasing volume of commodity imported.

7) The risk assessment should be amenable to updating when additional information becomes available.

Article 2.1.4.

Risk assessment steps

1. Entry assessment

Entry assessment consists of describing the biological pathway(s) necessary for an importation activity to introduce a pathogenic agent into a particular environment, and estimating the probability of that complete process occurring, either qualitatively (in words) or quantitatively (as a numerical estimate). The entry assessment describes the probability of the entry of each of the hazards (the pathogenic agents) or under each specified set of conditions
with respect to amounts and timing, and how these might change as a result of various actions, events or measures. Examples of the kind of inputs that may be required in the entry assessment are:

a) Biological factors
   - Species, strain or genotype, and age of aquatic animal
   - Strain of agent
   - Tissue sites of infection and/or contamination
   - Vaccination, testing, treatment and quarantine.

b) Country factors
   - Incidence or prevalence
   - Evaluation of Aquatic Animal Health Services, surveillance and control programmes, and zoning and compartmentalisation systems of the exporting country.

c) Commodity factors
   - Whether the commodity is alive or dead
   - Quantity of commodity to be imported
   - Ease of contamination
   - Effect of the various processing methods on the pathogenic agent in the commodity
   - Effect of storage and transport on the pathogenic agent in the commodity.

If the entry assessment demonstrates no significant risk, the risk assessment does not need to continue.

2. Exposure assessment

Exposure assessment consists of describing the biological pathway(s) necessary for exposure of animals and humans in the importing country to the hazards (in this case the pathogenic agents) from a given risk source, and estimating the probability of these exposure(s) occurring, either qualitatively (in words) or quantitatively (as a numerical estimate).

The probability of exposure to the identified hazards is estimated for specified exposure conditions with respect to amounts, timing, frequency, duration of exposure, routes of exposure, and the number, species and other characteristics of the animal and human populations exposed. Examples of the kind of inputs that may be required in the exposure assessment are:

a) Biological factors
   - Properties of the agent (e.g. virulence, pathogenicity and survival parameters)
   - Genotype of host.

b) Country factors
   - Presence of potential vectors or intermediate hosts
   - Aquatic animal demographics (e.g. presence of known susceptible species, distribution)
   - Human and terrestrial animal demographics (e.g. possibility of scavengers, presence of piscivorous birds)
   - Customs and cultural practices
   - Geographical and environmental characteristics (e.g. hydrographic data, temperature ranges, water courses).

c) Commodity factors
   - Whether the commodity is alive or dead
   - Quantity of commodity to be imported
   - Intended use of the imported aquatic animals or products (e.g. domestic consumption, restocking, incorporation in or use as aquaculture feed or bait)
   - Waste disposal practices.

If the exposure assessment demonstrates no significant risk, the risk assessment may conclude at this step.

3. Consequence assessment

Consequence assessment consists of describing the relationship between specified exposures to a biological agent and the consequences of those exposures. A causal process should exist by which exposures produce adverse health or environmental consequences, which may in turn lead to socio-economic consequences. The consequence assessment describes the potential consequences of a given exposure and estimates the probability
of them occurring. This estimate may be either qualitative (in words) or quantitative (a numerical estimate). Examples of consequences include:

a) Direct consequences
   - Aquatic animal infection, disease, production losses and facility closures
   - Public health consequences.

b) Indirect consequences
   - Surveillance and control costs
   - Compensation costs
   - Potential trade losses
   - Adverse, and possibly irreversible, consequences to the environment.

4. Risk estimation

Risk estimation consists of integrating the results of the entry assessment, exposure assessment, and consequence assessment to produce overall measures of risks associated with the hazards identified at the outset. Thus risk estimation takes into account the whole of the risk pathway from hazard identified to unwanted outcome.

For a quantitative assessment, the final outputs may include:
- The various populations of aquatic animals and/or estimated numbers of aquaculture establishments or people likely to experience health impacts of various degrees of severity over time
- Probability distributions, confidence intervals, and other means for expressing the uncertainties in these estimates
- Portrayal of the variance of all model inputs
- A sensitivity analysis to rank the inputs as to their contribution to the variance of the risk estimation output
- Analysis of the dependence and correlation between model inputs.

Article 2.1.5.

Principles of risk management

1) Risk management is the process of deciding upon and implementing measures to address the risks identified in the risk assessment, whilst at the same time ensuring that negative effects on trade are minimised. The objective is to manage risk appropriately to ensure that a balance is achieved between a country's desire to minimise the likelihood or frequency of disease incursions and their consequences and its desire to import commodities and fulfil its obligations under international trade agreements.

2) The international standards of the OIE are the preferred choice of sanitary measures for risk management. The application of these sanitary measures should be in accordance with the intentions of the standards.

Article 2.1.6.

Risk management components

1) Risk evaluation - the process of comparing the risk estimated in the risk assessment with the reduction in risk expected from the proposed risk management measures.

2) Option evaluation - the process of identifying, evaluating the efficacy and feasibility of, and selecting measures to reduce the risk associated with an importation. The efficacy is the degree to which an option reduces the likelihood or magnitude of adverse health and economic consequences. Evaluating the efficacy of the options selected is an iterative process that involves their incorporation into the risk assessment and then comparing the resulting level of risk with that considered acceptable. The evaluation for feasibility normally focuses on technical, operational and economic factors affecting the implementation of the risk management options.

3) Implementation - the process of following through with the risk management decision and ensuring that the risk management measures are in place.

4) Monitoring and review - the ongoing process by which the risk management measures are continuously audited to ensure that they are achieving the results intended.
Principles of risk communication

1) *Risk communication* is the process by which information and opinions regarding hazards and risks are gathered from potentially affected and interested parties during a *risk analysis*, and by which the results of the *risk assessment* and proposed *risk management* measures are communicated to the decision-makers and interested parties in the importing and exporting countries. It is a multidimensional and iterative process and should ideally begin at the start of the *risk analysis* process and continue throughout.

2) A *risk communication* strategy should be put in place at the start of each *risk analysis*.

3) The *communication of risk* should be an open, interactive, iterative and transparent exchange of information that may continue after the decision on importation.

4) The principal participants in *risk communication* include the authorities in the exporting country and other stakeholders such as domestic aquaculturists, recreational and commercial fishermen, conservation and wildlife groups, consumer groups, and domestic and foreign industry groups.

5) The assumptions and uncertainty in the model, model inputs and the *risk* estimates of the *risk assessment* should be communicated.

6) Peer review of *risk analyses* is an essential component of *risk communication* in order to obtain a scientific critique and to ensure that the data, information, methods and assumptions are the best available.

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NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2016.
SECTION 3.

QUALITY OF AQUATIC ANIMAL HEALTH SERVICES

CHAPTER 3.1.

QUALITY OF AQUATIC ANIMAL HEALTH SERVICES

Article 3.1.1.

The quality of Aquatic Animal Health Services depends on a set of factors, which include fundamental principles of an ethical, organisational, legislative, regulatory and technical nature. The Aquatic Animal Health Services shall conform to these fundamental principles, regardless of the political, economic or social situation of their country.

Compliance with these fundamental principles by a Member Country’s Aquatic Animal Health Service is important in the establishment and maintenance of confidence in its aquatic animal health status and international aquatic animal health certificates provided by the Aquatic Animal Health Service of other Member Countries.

These fundamental principles are presented in Article 3.1.2. Other factors to consider when evaluating Aquatic Animal Health Services are described in the Aquatic Code (notification, principles of certification, etc.).

The ability of Aquatic Animal Health Services to deliver appropriate services, monitor and control aquatic animal diseases based on Member Countries’ aquatic animal health legislation and regulations, can be measured through an evaluation or audit whose general principles are described in Articles 3.1.3. and 3.1.4.

A procedure for evaluating Aquatic Animal Health Services by OIE experts, on a voluntary basis, is described in Article 3.1.5.

Article 3.1.2.

Fundamental principles of quality

Aquatic Animal Health Services should comply with the following principles to ensure the quality of their activities:

1. Professional judgement
   Aquatic Animal Health Services should ensure that personnel have the relevant qualifications, scientific expertise and experience to give them the competence to make sound professional judgements.

2. Independence
   Care should be taken to ensure that the Aquatic Animal Health Service personnel are free from any commercial, financial, hierarchical, political or other pressures which may inappropriately influence their judgement or decisions.

3. Impartiality
   Aquatic Animal Health Services should be impartial. In particular, all the parties affected by their activities have a right to expect their services to be delivered under reasonable and non-discriminatory conditions.
Chapter 3.1.- Quality of Aquatic Animal Health Services

4. **Integrity**

Aquatic Animal Health Services are responsible for ensuring that the work of each of their personnel is of a consistently high level of integrity. Any fraud, corruption or falsification should be identified, documented and corrected.

5. **Objectivity**

Aquatic Animal Health Services should conduct themselves, in an objective, transparent and non-discriminatory manner.

6. **Aquatic animal health legislation and regulations**

Aquatic animal health legislation and regulations are a fundamental element that supports good governance and provides the legal framework for all key activities of the Aquatic Animal Health Service. Legislation and regulations should be suitably flexible to allow for judgements of equivalence and efficient responses to changing situations. In particular, they should define and document the responsibilities and structure of the organisations in charge of traceability and control of aquatic animal movements, aquatic animal disease control and reporting systems, epidemiological surveillance and communication of epidemiological information.

7. **General organisation**

Aquatic Animal Health Services should be able to demonstrate that they are able to anticipate the requirements for, and have control of, the establishment and application of aquatic animal health measures, and of international aquatic animal health certification activities. This should be demonstrated by means of appropriate legislation and regulations, sufficient financial resources and effective organisation.

Aquatic Animal Health Services should have at their disposal effective systems for aquatic animal disease surveillance, diagnosis and notification of disease problems that may occur in the national territory, in accordance with the provisions of the Aquatic Code. They should at all times endeavour to improve their performance in terms of aquatic animal health information systems and aquatic animal disease control.

Aquatic Animal Health Services should define and document the responsibilities and structure of the organisation (in particular the chain of command) in charge of issuing international aquatic animal health certificates.

Each position within the Aquatic Animal Health Services that has an impact on their quality should be described. These job descriptions should include the requirements for education, training, technical knowledge and experience.

8. **Quality policy**

Aquatic Animal Health Services should define and document their policy and objectives for, and commitment to, quality, and should ensure that this policy is understood, implemented and maintained at all levels in the organisation. Where conditions allow, they may implement a quality system corresponding to their areas of activity and appropriate for the type, range and volume of work that they have to perform. The recommendations provided in this chapter describe a suitable reference system, which should be used if a Member Country chooses to adopt a quality system.

9. **Procedures and standards**

Aquatic Animal Health Services should develop and document appropriate procedures and standards for all providers of relevant activities and associated facilities. These procedures and standards may for example relate to:

- a) programming and management of activities, including international aquatic animal health certification activities;
- b) prevention, control and notification of disease outbreaks;
- c) risk analysis, epidemiological surveillance and zoning;
- d) emergency preparedness for disasters which could have an impact on aquatic animal health and welfare of farmed fish;
- e) inspection and sampling techniques;
- f) diagnostic tests for aquatic animal diseases;
- g) preparation, production, registration and control of biological products for use in the diagnostic or prevention of diseases;
- h) border controls and import regulations;
- i) disinfection;
treatments intended to inactivate pathogens in aquatic animal products.

Where there are standards in the Aquatic Code or in the Aquatic Manual, Aquatic Animal Health Services should comply with these standards when applying aquatic animal health measures and when issuing international aquatic animal health certificates.

10. Information, complaints and appeals
Aquatic Animal Health Services should undertake to reply to requests from Aquatic Animal Health Services of other Member Countries or any other authority, in particular ensuring that any requests for information, complaints or appeals that are presented are dealt with in a timely manner.

A record should be maintained of all complaints and appeals and of the relevant action taken by Aquatic Animal Health Services.

11. Documentation
Aquatic Animal Health Services should have at their disposal a reliable and up-to-date documentation system suited to their activities.

12. Self-evaluation
Aquatic Animal Health Services should undertake periodical self-evaluation especially by documenting achievements against goals, and demonstrating the effectiveness of their organisational components and resource adequacy.

A procedure for evaluating Aquatic Animal Health Services by OIE experts, on a voluntary basis, is described in Article 3.1.5.

13. Communication
Aquatic Animal Health Services should have effective internal and external systems of communication covering administrative and technical staff and parties affected by their activities.

14. Human and financial resources
Responsible authorities should ensure that adequate resources are made available to implement effectively the above activities.

Article 3.1.3.

For the purposes of the Aquatic Code, every Member Country should recognise the right of another Member Country to undertake, or request it to undertake, an evaluation of its Aquatic Animal Health Services where the initiating Member Country is an actual or a prospective importer of aquatic animal commodities and/or where the evaluation is to be a component of a risk analysis process that is to be used to determine or review sanitary measures which apply to such trade.

A Member Country has the right to expect that the evaluation of its Aquatic Animal Health Services will be conducted in an objective and transparent manner. A Member Country undertaking an evaluation should be able to justify any measure taken as a consequence of its evaluation.

Article 3.1.4.

A Member Country which intends to conduct an evaluation of another Member Country's Aquatic Animal Health Services should provide notice in writing, and allow sufficient time for the other Member Country to comply with the request. This notice should define the purpose of the evaluation and details of the information required.

On receipt of a formal request for information to enable an evaluation of its Aquatic Animal Health Services by another Member Country, and following bilateral agreement of the evaluation process and criteria, a Member Country should expeditiously provide the Member Country requesting the evaluation with meaningful and accurate information of the type requested.

The evaluation process should take into account the fundamental principles and other factors of quality laid down in Article 3.1.1. and in Article 3.1.2. It should also take into consideration the specific circumstances regarding quality, as described in Article 3.1.1., prevailing in the countries concerned.
Chapter 3.1.- Quality of Aquatic Animal Health Services

The outcome of an evaluation conducted by a Member Country should be provided in writing as soon as possible, and in any case within four months of receipt of the relevant information, to the Member Country which has undergone the evaluation. The evaluation report should detail any findings that affect trade prospects. The Member Country which conducts the evaluation should clarify in detail any points of the evaluation on request.

In the event of a dispute between two Member Countries over the conduct or the conclusions of the evaluation of Aquatic Animal Health Services, the matter should be dealt with having regard to the procedures set out in Article 3.1.3.

Article 3.1.5.

Evaluation facilitated by OIE experts under the auspices of the OIE

The OIE has established procedures for the evaluation of Aquatic Animal Health Services of Member Countries. Member Countries can make a request to the OIE for an evaluation of their Aquatic Animal Health Services.

The World Assembly of OIE Delegates may endorse a list of approved experts to facilitate the evaluation process.

Under these procedures, the Director General of the OIE recommends an expert(s) from that list.

The expert(s) facilitate(s) the evaluation of the Aquatic Animal Health Services of the Member Country using the OIE Performance of Veterinary Services and/or Aquatic Animal Health Services (OIE PVS Tool: Aquatic).

The expert(s) produce(s) a report in consultation with the Veterinary Services of the Member Country.

The report is submitted to the Director General of the OIE and, with the consent of the Member Country, published by the OIE.

NB: FIRST ADOPTED IN 2009; MOST RECENT UPDATE ADOPTED IN 2014.
CHAPTER 3.2.

COMMUNICATION

Article 3.2.1.

General considerations

In general, communication entails the exchange of information between various individual, institutional and public groups for purposes of informing, guiding and motivating action. The application of the science and technique of communication involves modulating messages in accordance with situations, objectives and target audiences.

The recognition of communication as a discipline of the Aquatic Animal Health Services and its incorporation within it is critical for their operations. The integration of aquatic animal health and communication expertises is essential for effective communication. Communication between the Aquatic Animal Health Services and Veterinary Services (particularly where Aquatic Animal Health Services are separate from, and independent of Veterinary Services) is especially important.

Communication should be an integral part of all the activities of the Aquatic Animal Health Services including animal health (surveillance, early detection and rapid response, prevention and control), aquatic animal welfare and veterinary public health (food safety, zoonoses) and veterinary medicine.

Objectives of this chapter on communication for the Aquatic Animal Health Services are to provide guidance for the development of a communication system, strategic and operational communication plans and elements to assess their quality.

Article 3.2.2.

Principles of communication

1) Aquatic Animal Health Services should have the authority and capability to communicate on matters within their mandate.

2) Aquatic animal health and communication expertises should be combined.

3) Communication should be targeted and follow the fundamental criteria of transparency, consistency, timeliness, balance, accuracy, honesty and empathy and respect the fundamental principles of quality of Aquatic Animal Health Services (Article 3.1.2.)

4) Communication should be a continuous process.

5) Aquatic Animal Health Services should have oversight of planning, implementing, monitoring, evaluating and revising their strategic and operational communication plans.

Article 3.2.3.

Definitions

Communication: means the discipline of informing, guiding and motivating individual, institutional and public groups, ideally on the basis of interactive exchanges, about any issue under the competence of the Aquatic Animal Health Services.

Crisis: means a situation of great threat, difficulty or uncertainty when issues under the competence of the Aquatic Animal Health Services require immediate action.

Crisis communication: means the process of communicating information as accurately as possible, albeit potentially incomplete, within time constraints in the event of a crisis.

Outbreak communication: means the process of communicating in the event of an outbreak. Outbreak communication includes notification.
Article 3.2.4.

Communication system

In addition to the Principles of Communication the following elements should be used in conjunction with Chapter 3.1., when planning, implementing and assessing a communication system:

1. **Organisational chart** indicating a direct link between the communication personnel and the Competent Authority, through the chain of command such as dedicated communication unit and communication officer

2. **Human resources**
   a) Identified and accessible official communication focal point
   b) Job descriptions of communication personnel identifying roles and responsibilities
   c) Sufficient number of qualified personnel with knowledge, skills, attitude and abilities relevant to communication
   d) Continuous training and education on communication provided to communication personnel.

3. **Financial and physical resources**
   a) Clearly identified budget for communication that provides adequate funding
   b) Provision or access to appropriate material resources in order to carry out roles and responsibilities: suitable premises or accommodation that is adequately equipped with sufficient office and technical equipment, including information technology and access to the Internet.

4. **Management of the communication system**
   a) Roles and responsibilities of the communication personnel
      i) Report to the Competent Authority
      ii) Engage in decision-making process by providing guidance and expertise on communication issues to the Competent Authority
      iii) Be responsible for the planning, implementation and evaluation of the strategic and operational plans for communication and relevant standard operating procedures
      iv) Function as contact point on communication issues for the Aquatic Animal Health Services
      v) Provide and coordinate continuous education on communication for the Aquatic Animal Health Services.
   b) **Strategic plan for communication**

      A well-designed strategic plan for communication should support the Aquatic Animal Health Services strategic plan and have management support and commitment. The strategic plan for communication should address all high level organization-wide long-term communication objectives.

      A strategic plan for communication should be monitored and periodically reviewed, and should identify measurable performance objectives and techniques to assess the effectiveness of communication.

      The strategic plan for communication should consider the different types of communication: routine communication, risk communication, outbreak communication and crisis communication, to allow individuals, affected or interested parties, an entire community or the general public to make the best possible decisions and be informed of policy decisions and their rationale.

      The key outcomes in effectively implementing a strategic plan for communication are increased knowledge and awareness of issues by the public and stakeholders, higher understanding of the role of the Aquatic Animal Health Services, higher visibility of and improved trust and credibility in the Aquatic Animal Health Services. These will enhance understanding and/or acceptance of policy decisions and subsequent change of perception, attitude and/or behaviour.
c) Operational plans for communication

Operational plans for communication should be based on the assessment of specific issues and should identify specific objectives and target audiences such as staff, partners, stakeholders, media and the general public.

Each operational plan for communication should consist of a well-planned series of activities using different techniques, tools, messages and channels to achieve intended objectives and utilising available resources within a specific timeframe.

NB: FIRST ADOPTED IN 2012.
SECTION 4.
DISEASE PREVENTION AND CONTROL

CHAPTER 4.1.
ZONING AND COMPARTMENTALISATION

Article 4.1.1.

Introduction

Given the difficulty of establishing and maintaining freedom from a particular disease for an entire country especially for diseases whose entry is difficult to control, there may be benefits to one or more Member Countries in establishing and maintaining a subpopulation with a distinct aquatic animal health status. Subpopulations may be separated by natural or artificial geographical barriers or, in certain situations, by the application of appropriate management practices.

Zoning and compartmentalisation are procedures implemented by a country under the provisions of this chapter to define subpopulations of distinct aquatic animal health status for the purpose of disease control or international trade. Compartmentalisation applies to a subpopulation when management practices related to biosecurity are the defining factors, while zoning applies when a subpopulation is defined on a geographical basis. In practice, spatial considerations and good management play important roles in the application of both concepts.

This chapter is to assist Member Countries wishing to establish and maintain different subpopulations, using the principles of compartmentalisation and zoning. These principles should be applied in accordance with the measures recommended in the relevant disease chapter(s). This chapter also outlines a process through which trading partners may recognise such subpopulations. This process is best implemented by trading partners through establishing parameters and gaining agreement on the necessary measures prior to outbreaks of disease.

Before trade in aquatic animals or aquatic animal products may occur, an importing country needs to be satisfied that its aquatic animal health status will be appropriately protected. In most cases, the import regulations developed will rely in part on judgements made about the effectiveness of sanitary procedures undertaken by the exporting country, both at its borders and within its territory.

In addition to contributing to the safety of international trade, zoning and compartmentalisation may assist disease control or eradication within Member Countries. Zoning may encourage the more efficient use of resources, and compartmentalisation may allow the functional separation of a subpopulation from other domestic or wild aquatic animals through biosecurity measures, which a zone (through geographical separation) would not achieve. Following an outbreak of disease, compartmentalisation may allow a Member Country be able to take advantage of epidemiological links among subpopulations or common practices relating to biosecurity, despite diverse geographical locations, to facilitate disease control and/or the resumption of trade.

Zoning and compartmentalisation may not be applicable to all diseases, but separate requirements will be developed for each disease for which the application of zoning or compartmentalisation is considered appropriate.

To regain the status of a free zone or free compartment following an outbreak of disease, Member Countries should follow the recommendations in the relevant disease chapter in the Aquatic Code.
Article 4.1.2.

General considerations

The Competent Authority of an exporting country that is establishing a zone or compartment for international trade purposes should clearly define the subpopulation in accordance with the recommendations in the relevant chapters in the Aquatic Code, including those on surveillance, and the identification and traceability of aquatic animals. The Competent Authority of an exporting country should be able to explain to the Competent Authority of an importing country the basis for its claim of a distinct aquatic animal health status for the zone or compartment in such terms.

The procedures used to establish and maintain the distinct aquatic animal health status of a zone or compartment should be appropriate to the particular circumstances and will depend on the epidemiology of the disease, environmental factors, risk of introduction and establishment of disease, and applicable biosecurity measures. The exporting country should be able to demonstrate, through detailed documentation supplied to the importing country, published through official channels, that it has implemented the recommendations in the Aquatic Code for establishing and maintaining such a zone or compartment.

An importing country should recognise the existence of this zone or compartment when the appropriate measures recommended in the Aquatic Code are applied, and the Competent Authority of the exporting country certifies that this is the case. Note that an importing country may adopt a higher level of protection where it is scientifically justified and the obligations referred to in Article 5.3.1 are met.

Where countries share a zone or compartment, the Competent Authority of each country should cooperate to define and fulfill their respective responsibilities.

The exporting country should conduct an assessment of the resources needed and available to establish and maintain a zone or compartment for international trade purposes. These include the human and financial resources and the technical capability of the Aquatic Animal Health Service (and of the relevant industry, in the case of a compartment) including disease surveillance and diagnosis.

Article 4.1.3.

Principles for defining a zone or compartment, including protection zones

In conjunction with the above considerations and the definitions of zone and compartment, the following principles should apply when Member Countries define a zone or compartment:

1) The extent of a zone should be established by the Aquatic Animal Health Service on the basis of the definition of zone and made public through official channels.

2) A protection zone may be established to preserve the health status of aquatic animals in a free country or free zone, from adjacent countries or zones of different aquatic animal health status. Measures should be implemented based on the epidemiology of the disease under consideration to prevent introduction of the pathogenic agent. These measures should include intensified movement control and surveillance and may include vaccination, raised awareness or other measures. The application of these measures can be in the entire free zone or in a defined area within and/or outside the free zone.

3) The factors defining a compartment should be established by the Aquatic Animal Health Service on the basis of relevant criteria such as management and husbandry practices related to biosecurity, and made public through official channels.

4) Aquatic animals belonging to such subpopulations need to be recognisable as such through a clear epidemiological separation from other aquatic animals and all things presenting a disease risk.

5) For a zone or compartment, the Aquatic Animal Health Service should document in detail the measures taken to ensure the identification of the subpopulation, for example by means of registration of all the aquaculture establishments located in such a zone or compartment and the establishment and maintenance of its aquatic animal health status through a biosecurity plan. The measures used to establish and maintain the distinct aquatic animal health status of a zone or compartment should be appropriate to the particular circumstances and will depend on the epidemiology of the disease, environmental factors, the aquatic animal health status in adjacent areas, applicable biosecurity measures (including movement controls, use of natural and artificial boundaries, the spatial separation of aquatic animals, and commercial management and husbandry practices), and surveillance.

6) For a compartment, the biosecurity plan should describe the partnership between the relevant enterprise/industry and the Aquatic Animal Health Service, and their respective responsibilities, including the procedures for oversight of the operation of the compartment by the Aquatic Animal Health Service.
7) For a compartment, the biosecurity plan should also describe the routine operating procedures to provide clear evidence that the surveillance conducted and the management practices are adequate to meet the definition of the compartment. In addition to information on aquatic animal movements, the biosecurity plan should include production and stock records, feed sources, traceability, surveillance results, visitor logbook, morbidity and mortality history, medications, vaccinations, water supply and effluent treatments, documentation of training and any other criteria necessary for evaluation of risk mitigation. The information required may vary in accordance with the aquatic animal species and disease(s) under consideration. The biosecurity plan should also describe how the measures will be audited to ensure that the risks are regularly re-assessed and the measures adjusted accordingly.

8) Thus defined, the zones and compartments constitute the relevant subpopulations for the application of the recommendations in Sections 8 to 11.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2010.
CHAPTER 4.2.

APPLICATION OF COMPARTMENTALISATION

Article 4.2.1.

Introduction and objectives

The recommendations in this chapter provide a structured framework for the application and recognition of compartments within countries or zones, based on the provisions of Chapter 4.1. with the objective to facilitate trade in aquatic animals and products of aquatic animal origin and as a tool for disease management.

Establishing and maintaining a disease-free status throughout the country should be the ultimate goal for Member Countries. However, establishing and maintaining a disease-free status for an entire country may be difficult, especially in the case of diseases that exist in wild aquatic animal species or can easily cross international boundaries. For many diseases, Member Countries have traditionally applied the concept of zoning to establish and maintain an animal subpopulation with a different animal health status within national boundaries.

The essential difference between zoning and compartmentalisation is that the recognition of zones is based on geographical boundaries whereas the recognition of compartments is based on management and biosecurity practices. However, spatial considerations and good management practices play a role in the application of both concepts.

The fundamental requirement for compartmentalisation is the implementation and documentation of management and biosecurity measures to create a functional separation of subpopulations.

For example, an aquaculture establishment in an infected country or infected zone might have biosecurity measures and management practices that result in negligible risk from diseases or agents. The concept of a compartment extends the application of a ‘risk boundary’ beyond that of a geographical interface and considers all epidemiological factors that can help to create an effective disease-specific separation between subpopulations.

In disease free countries or free zones, it is preferable that compartments are defined prior to the occurrence of a disease outbreak. In the event of an outbreak or in infected countries or infected zones, compartmentalisation may be used to facilitate trade.

For the purpose of international trade, compartments should be under the responsibility of the Competent Authority in the country. For the purposes of this chapter, compliance by the Member Countries with Chapters 1.1. and 3.1. is an essential prerequisite.

Article 4.2.2.

Principles for defining a compartment

A compartment may be established with respect to a specific disease or diseases. A compartment should be clearly defined. This should indicate, inter alia, the location of all its components including establishments, as well as related functional units (such as brood stock facilities, hatcheries, nurseries, grow-out facilities, slaughterhouses, processing plants, etc.). It should also describe their interrelationships and their contribution to an epidemiological separation between the aquatic animals in a compartment and subpopulations elsewhere with a different health status. The definition of compartment should encompass disease-specific epidemiological factors, the aquatic animal species in the compartment, production systems, biosecurity practices, infrastructural factors and surveillance.

Article 4.2.3.

Separation of a compartment from potential sources of infection

The management of a compartment should provide to the Aquatic Animal Health Service documented evidence on the following:
1. Physical or spatial factors that affect the status of biosecurity in a compartment

While a compartment is primarily based on management and biosecurity measures, a review of geographical factors is needed to ensure that the functional boundary provides adequate separation of a compartment from adjacent animal populations with a different health status. The following factors should be taken into consideration in conjunction with biosecurity measures and, in some instances, may alter the degree of confidence achieved by general biosecurity and surveillance measures:

a) disease status in adjacent areas and in areas epidemiologically linked to the compartment;

b) location, disease status and biosecurity of the nearest epidemiological units or other epidemiologically relevant premises. Consideration should be given to the distance and physical separation from:

i) aquatic animal populations with a different health status in close proximity to the compartment, including wildlife and their migratory routes;

ii) slaughterhouses or processing plants;

iii) exhibitions, ‘put and take’ fisheries, fish markets, restaurants with live fish and other points of aquatic animal concentration.

2. Infrastructural factors

Structural aspects of an establishment or establishments within a compartment contribute to the effectiveness of its biosecurity. Consideration should be given to:

a) water supply;

b) effective means of physical separation;

c) facilities for people entry including access control;

d) vehicle and vessel access including washing and disinfection procedures;

e) unloading and loading facilities;

f) isolation facilities for introduced aquatic animals;

g) facilities for the introduction of material and equipment;

h) infrastructure to store feed and veterinary products;

i) disposal of aquatic animal waste;

j) measures to prevent exposure to fomites or vectors;

k) feed supply/source.

3. Biosecurity plan

The integrity of the compartment relies on effective biosecurity. The management of the compartment should develop, implement and monitor a comprehensive biosecurity plan.

The biosecurity plan should describe in detail:

a) potential pathways for introduction and spread into the compartment of the agents for which the compartment was defined, including aquatic animal movements, wild aquatic animals, potential vectors, vehicles, people, biological products, equipment, fomites, feed, waterways, drainage or other means. Consideration should also be given to the survivability of the agent in the environment;

b) the critical control points for each pathway;

c) measures to mitigate exposure for each critical control point;

d) standard operating procedures including:

i) implementation, maintenance, monitoring of compliance with the risk mitigation measures;

ii) application of corrective actions;

iii) verification of the process;

iv) record keeping;

e) contingency plan in the event of a change in the level of exposure;

f) reporting procedures to the Competent Authority;

g) the programme for educating and training workers to ensure that all persons involved are knowledgeable and informed on biosecurity principles and practices;

h) the surveillance programme in place.

In any case, sufficient evidence should be submitted to assess the efficacy of the biosecurity plan in accordance with the level of risk for each identified pathway. This evidence should be structured in line with the principles of Hazard Analysis and Critical Control Point (HACCP). The biosecurity risk of all operations of the compartment should be re-assessed and documented at least on a yearly basis. Based on the outcome of the assessment,
concrete and documented mitigation steps should be taken to reduce the likelihood of introduction of the pathogenic agent into the compartment.

4. Traceability system

A prerequisite for assessing the integrity of a compartment is the existence of a valid traceability system. Although individual identification of aquatic animals may not be feasible, the Competent Authority should provide sufficient assurance of traceability in such a way that their history and movements can be documented and audited.

All aquatic animal movements into and out of the compartment should be recorded at the compartment level, and when needed, based on a risk assessment, approved by the Competent Authority. Movements within the compartment need not be certified but should be recorded and documented at the compartment level.

Article 4.2.4.

Documentation

Documentation should provide clear evidence that the biosecurity, surveillance, traceability and management practices defined for a compartment are effectively and consistently applied. In addition to animal movement information, the necessary documentation should include production unit records (e.g. cage, pond), feed sources, laboratory tests, mortality records, visitor logbook, morbidity history, water supply and effluent treatments, medication and vaccination records, biosecurity plans, training documentation and any other criteria necessary for the evaluation of disease exclusion.

The historical status of a compartment for the disease(s) for which it was defined should be documented and demonstrate compliance with the requirements for freedom in the relevant chapter of the Aquatic Code.

In addition, a compartment seeking recognition should submit to the Competent Authority a baseline aquatic animal health report indicating the presence or absence of listed diseases. This report should be regularly updated to reflect the current aquatic animal health status of the compartment.

Vaccination records including the aquatic animal groups vaccinated, type of vaccine and frequency of administration should be available to enable interpretation of surveillance data.

The time period for which all records should be kept may vary in accordance with the species and disease(s) for which the compartment was defined.

All relevant information should be recorded in a transparent manner and be easily accessible so as to be auditable by the Competent Authority.

Article 4.2.5.

Surveillance for the pathogenic agent or disease

The surveillance system should comply with Chapter 1.4 on surveillance and the specific recommendations for surveillance for the disease(s) for which the compartment was defined, if available.

If there is an increased risk of exposure to the agent for which the compartment has been defined, the sensitivity of the internal and external surveillance system should be reviewed, documented and, where necessary, increased. At the same time, biosecurity measures in place should be reassessed and increased if necessary.

1. Internal surveillance

Surveillance should involve the collection and analysis of disease/infection data so that the Competent Authority can certify that the animal subpopulation contained in all the establishments comply with the defined status of that compartment. A surveillance system that is able to ensure early detection in the event that the agent enters a subpopulation is essential. Depending on the disease(s) for which the compartment was defined, different surveillance strategies may be applied to achieve the desired confidence in disease freedom.
2. **External surveillance**  
The biosecurity measures applied in a *compartment* should be appropriate to the level of exposure of the *compartment*. External surveillance will help identify a significant change in the level of exposure for the identified pathways for disease introduction into the *compartment*.  
An appropriate combination of targeted and passive surveillance is necessary to achieve the goals described above. Based on the recommendations of Chapter 1.4., targeted surveillance based on an assessment of risk factors may be the most efficient surveillance approach. Targeted surveillance should in particular include epidemiological units in close proximity to the *compartment* or those that have a potential epidemiological link with it.

**Article 4.2.6.**  
**Diagnostic capabilities and procedures**  
Officially-designated laboratory facilities should be available for sample testing. All laboratory tests and procedures should comply with the recommendations of the Aquatic Manual for the specific disease. Each laboratory that conducts testing should have systematic procedures in place for rapid reporting of disease results to the Competent Authority. Where appropriate, results should be confirmed by an OIE Reference Laboratory.

**Article 4.2.7.**  
**Emergency response and notification**  
Early detection, *diagnosis*, *notification* of disease and rapid response are critical to minimise the consequences of outbreaks.  
In the event of suspicion of occurrence of the disease for which the *compartment* was defined, the free status of the *compartment* should be immediately suspended. If confirmed, the status of the *compartment* should be immediately revoked and importing countries should be notified following the provisions of Chapter 1.1.  
In case of the detection of any disease not present in accordance with the baseline animal health report of the *compartment* referred to in Article 4.2.4., the management of the *compartment* should notify the Competent Authority, and initiate a review to determine whether there has been a breach in the biosecurity measures and notify the Competent Authority. If a significant breach in biosecurity, even in the absence of outbreak, is detected, export certification as a *free* compartment should be suspended. Disease-free status of the *compartment* may only be reinstated after the *compartment* has adopted the necessary measures to re-establish the original biosecurity level and the Competent Authority re-approves the status of the *compartment*.  
In the event of a *compartment* being at risk from a change, in the surrounding area, in the disease situation for which the *compartment* was defined, the Competent Authority should re-evaluate without delay the status of the *compartment* and consider whether any additional biosecurity measures are needed to ensure that the integrity of the *compartment* is maintained.

**Article 4.2.8.**  
**Supervision and control of a compartment**  
The authority, organisation, and infrastructure of the Aquatic Animal Health Services, including laboratories, should be clearly documented in accordance with Chapter 3.1., to provide confidence in the integrity of the *compartment*.  
The Competent Authority has the final authority in granting, suspending and revoking the status of a *compartment*. The Competent Authority should continuously supervise compliance with all the requirements critical to the maintenance of the *compartment* status described in this chapter and ensure that all the information is readily accessible to the importing countries. Any significant change should be notified to the importing country.

NB: FIRST ADOPTED IN 2010; MOST RECENT UPDATE ADOPTED IN 2016.
CHAPTER 4.3.

DISINFECTION OF AQUACULTURE ESTABLISHMENTS AND EQUIPMENT

Article 4.3.1.

Purpose

To provide recommendations on planning and implementation of disinfection procedures to prevent the introduction, establishment or spread of pathogenic agents.

Article 4.3.2.

Scope

This chapter describes recommendations for disinfection of aquaculture establishments and equipment during routine biosecurity activities and for emergency response. Guidance is provided on general principles, planning and implementation of disinfection activities.

For specific methods of pathogen inactivation refer to the disease-specific chapters in the Aquatic Manual.

Article 4.3.3.

Introduction

Disinfection is employed as a disease management tool in aquaculture establishments as part of a biosecurity plan. Disinfection is used to prevent entry or exit of target pathogenic agents to or from an aquaculture establishment or compartment, as well as the spread of pathogenic agents within aquaculture establishments. Disinfection may be used during emergency disease response to support the maintenance of disease control zones and for disease eradication (stamping-out procedures) from affected aquaculture establishments. The specific objective of disinfection will determine the strategy used and how it is applied.

When possible, the spread of pathogenic agents should be prevented by avoiding transmission pathways rather than attempting to manage them through disinfection. For example, difficult to disinfect items (e.g. gloves, dive and harvest equipment, ropes and nets) should be dedicated to a specific site rather than moved between production units or aquaculture establishments after disinfection.

Article 4.3.4.

General principles

Disinfection is a structured process that uses physical and chemical procedures to remove organic material and destroy or inactivate pathogenic agents. The process should include planning and implementation stages that take into account potential options, efficacy and risks.

The disinfection process may vary depending on whether the overall objective is disease prevention, control or eradication. Procedures addressing eradication will generally involve destocking of all aquatic animals as well as disinfection of aquaculture establishments and equipment, whereas disease control aims at limiting the spread of
Chapter 4.3.- Disinfection of aquaculture establishments and equipment

disease between or within aquaculture establishments. Although different approaches may be used to achieve the identified objective, the general principles described below should be applied in all cases.

1) The disinfection process should include the following phases:
   a) Cleaning and washing
   Cleaning and washing of surfaces and equipment is necessary to remove solid waste, organic matter (including biofouling) and chemical residues as these may reduce the efficacy of disinfectants. The use of detergent is also important to break down biofilms. The detergent used should be compatible with the disinfectant and the surface being treated. After cleaning, any excess water should be drained and before the application of disinfectants all surfaces and equipment should be inspected to ensure there is no remaining organic material.
   Where treatment of water is required, the presence of suspended solids may also reduce the efficacy of some disinfectants. Removal of suspended solids through various processes such as filtration, sedimentation, coagulation or flocculation should be performed.
   Biofilms, often referred to as slime, are a thin film of microorganisms and extracellular polymeric substances that adhere to surfaces. Biofilms physically protect embedded microorganisms against disinfectants. In order to achieve effective disinfection, biofilms should be removed during the cleaning and washing stage prior to the application of disinfectants.
   All waste produced should be disposed of in a biosecure manner because it may contain viable pathogenic agents that have the potential to spread infection if not controlled.

b) Application of disinfectants
   This phase involves the application of chemical compounds or physical processes that are appropriate to inactivate the pathogenic agent.
   The application of disinfectants should take into account the type of material requiring disinfection and how disinfectants should be applied. Hard non-permeable materials (e.g. polished metal surfaces, plastics and painted concrete) can be cleaned thoroughly and allow contact with the disinfectant because there is little opportunity for infective material to lodge in crevices. Disinfection efficacy will decrease if the surface is corroded, pitted or paint is flaking, therefore proper maintenance of surfaces and equipment is essential. For permeable surfaces and materials (e.g. woven material, nets and soil), a higher disinfectant concentration and a longer contact time is required because the surface area is greater, chemicals cannot penetrate easily and residual organic matter may be present.
   The choice of the application method should ensure all surfaces come into contact with the agent for the required period of time. The application of disinfectants should be undertaken methodically (e.g. using a grid pattern) to ensure that complete coverage and adequate contact times are achieved. Each phase should start from the highest point and proceed downwards, commencing from the least contaminated areas. However for some equipment, rinsing of surfaces with the disinfectant may be sufficient. When disinfectants are applied to vertical surfaces, care should be taken to ensure that the required contact time is maintained before the disinfectant drains away. Vertical surfaces may need retreatment or require the addition of compatible foaming agents to prolong adherence to surfaces.
   For pipes and biofilters, complete filling with the disinfectant solution should be done to ensure contact with all surfaces. Difficult to access and complex areas may require fumigation or use of misting equipment.

   c) Removal or inactivation of the disinfectant
   Removal or inactivation of chemical residues is important to avoid toxicity to aquatic animals, corrosion of equipment and environmental impacts. Processes that may be employed for the removal or inactivation of chemical residues may include: rinsing of surfaces, dilution to acceptable levels, treatment to inactivate chemical agents or, time to allow deactivation or dissipation of the active compound. These processes may be used in isolation or in combination.

   2) Disinfectants should be used in accordance with relevant legislation. Disinfectants may present risks to the health of people, aquatic animals and the environment. Chemical disinfectants should be stored, used and disposed of in accordance with regulations and manufacturer's instructions.

   3) Disinfection should be monitored to ensure appropriate dose of disinfectant and disinfection efficacy. Depending on the application process and the pathogenic agent of concern, this may be done in different ways. Examples include measurement of the active agent (e.g. residual chlorine levels), indirect measurement of the active agent by an indicator process (e.g. monitoring oxygen reduction potential), and measuring efficacy using indicator bacteria (e.g. heterotrophic bacteria plate counts).
   In facilities that have undergone destocking and disinfection, the use of a sentinel population prior to restocking may be considered. The sentinel population should be susceptible to the pathogen of concern and exposed to conditions that would be conducive to the expression of clinical disease should viable pathogen remain.

   4) Aquaculture establishments should keep records of the disinfection processes applied. The records should be sufficient to allow evaluation of the disinfection plan.
Planning

A disinfection plan should be developed that incorporates an assessment of the transmission pathways, the type of material to be disinfected, the pathogenic agents to be inactivated, the health and safety precautions and control measures required, and the environment in which the process is to be undertaken. The disinfection plan should include a mechanism for determining efficacy. The disinfection plan should be regularly reviewed to ensure the disinfection process remains effective and efficient. Any changes to the disinfection plan should also be documented.

The planning process should assess the critical control points where disinfection will be most effective. Disinfection priorities should be developed by considering potential pathways for spread of pathogenic agents and the relative likelihood of contamination. For effective disinfection of facilities containing vectors (e.g. ponds) the vectors should be excluded, removed or destroyed as part of the disinfection process.

An inventory of all items requiring disinfection should be developed when practical. An assessment should be made of the materials used in construction, their surface porosity and resistance to chemical damage, and accessibility for disinfection. Then, the appropriate disinfection method should be decided for each item.

The level of cleaning required prior to disinfection should be assessed for each type of equipment. If heavy soiling with solids and particulate matter is present, specific attention should be given to the cleaning process and the resources required. The physical or chemical cleaning process should be compatible with the disinfectant chosen.

Personnel, equipment and materials to be disinfected should be assessed taking into account the type and number of items to be treated and how waste material will be managed.

The ability to control water flow and water volumes should be considered at the planning stage and will depend on the enterprise type (recirculation, flow-through and open systems). Water may be disinfected using a variety of methods as described in Article 4.3.11.

Disinfection in an emergency response

Disinfection is an essential part of any emergency response to support disease control activities such as quarantine of affected aquaculture establishments and stamping-out procedures. The conditions associated with an emergency response require different approaches for disinfection to those used in routine biosecurity. These conditions include a high level of disease risk (due to the significance of the disease), high pathogen loading, potential high volumes of infected aquatic animals and waste, large areas requiring disinfection and large volumes of contaminated water. Planning should consider these circumstances, incorporate an evaluation of risks and include methods for monitoring efficacy.

In an emergency response it may be preferable to avoid transmission pathways rather than relying on disinfection. Equipment should not be moved from an infected aquaculture establishment unless effective disinfection has been achieved. In some circumstances, equipment or material that is difficult to disinfect or has a high likelihood of contamination may need to be disposed of in a biosecure manner rather than be disinfected.

Types of disinfectants

Types of disinfectants commonly used in aquaculture include the following:

1. Oxidising agents
   The majority of oxidising agents are relatively fast acting and are effective disinfectants for a large range of micro-organisms. These compounds are inactivated by organic matter and therefore should be used following an effective cleaning stage. Organic matter consumes oxidising agents and the initial concentration (loading dose) may drop rapidly, making effective dosing levels (residual dose) difficult to predict. Therefore, residual dose levels...
should always be monitored to ensure that they remain above the minimum effective concentration for the required time period.

Oxidising agents may be toxic to aquatic animals and therefore should be removed or inactivated. Common oxidising agents include chlorine compounds, chloramine-T, iodophores, peroxygen compounds, chlorine dioxide and ozone.

2. pH modifiers (alkalis and acids)

Modification of pH can be achieved through the use of either alkaline or acidic compounds. Advantages of using pH modifiers include the ease of determining their concentrations and that they are not inactivated by organic matter. Also, they can be used in areas where the application of other effective disinfectants is not possible, such as in pipes or on biofilter surfaces.

3. Aldehydes

Aldehydes act by denaturing protein. Two aldehyde compounds that may be used during decontamination of aquaculture establishments are formaldehyde and glutaraldehyde. They are highly effective against a wide range of organisms but require long exposure times. Aldehydes maintain their activity in the presence of organic matter and are only mildly corrosive. Glutaraldehyde is used in the liquid form as a cold sterilant, particularly for heat-sensitive equipment. Formaldehyde may be used as a mist or a gas for fumigation.

4. Biguanides

Of the many biguanides available, chlorhexidine is the most commonly used. However they are not effective in hard or alkaline water and are less effective against many pathogenic agents compared to other groups of disinfectants. These compounds are comparatively non-corrosive and relatively safe, thus they are commonly used in the disinfection of skin surfaces and delicate equipment.

5. Quaternary ammonium compounds (QACs)

The biocidal efficacy of QACs is variable and selective. They are effective against some vegetative bacteria and some fungi, but not all viruses. QACs are most active against gram-positive bacteria; action against gram-negative bacteria is slow, with some strains showing resistance. These compounds are not effective against spores. The advantages of QACs are that they are noncorrosive and have wetting properties that enhance contact with surfaces. QACs may be toxic to aquatic animals and should be removed from surfaces following disinfection procedures.

6. Ultraviolet (UV) irradiation

UV irradiation is a viable option for the treatment of water entering or leaving aquaculture establishments where there is some control of water flows in recirculation or flow-through systems. UV irradiation should be used following effective filtration because suspended solids reduce UV transmission and the effectiveness of this method.

7. Heat treatment

Susceptibility of pathogenic agents to heat treatment varies significantly. Under most conditions, moist heat is more effective than dry heat.

8. Desiccation

Desiccation may be an effective disinfectant for susceptible pathogenic agents and may be used in circumstances where other disinfection methods are impractical or as an ancillary method to other disinfection methods. Desiccation can be considered to be a disinfection method if complete drying of the item is achieved because the absence of water will kill many pathogenic agents. However, moisture content may be difficult to monitor in some circumstances. The effectiveness will vary depending on environmental conditions such as temperature and humidity.

9. Combined disinfection methods

Combined disinfection methods should be considered wherever they are synergistic and provide a higher assurance of effective pathogenic agent inactivation. Some examples include:

   a) direct sunlight and desiccation as a combined disinfection method provides three potential disinfection actions, i.e. UV irradiation, heating and desiccation. It has no operational cost and may be used subsequent to other methods;
b) Ozone and UV irradiation are often combined in series as they provide back-up systems and different modes of action. UV irradiation also has the advantage of removing ozone residues from treated water. Antagonistic effects may occur when chemical agents or detergents are combined.

Article 4.3.8.

Selection of a disinfectant

The disinfectant should be selected considering the following:

- efficacy against the pathogenic agents;
- effective concentration and exposure time;
- ability to measure efficacy;
- nature of the items to be disinfected and the potential for them to be damaged;
- compatibility with the available water type (e.g. fresh water, hard water or seawater);
- availability of the disinfectant and equipment;
- ease of application;
- the ability to remove organic matter;
- cost;
- impacts of residues on aquatic animals and the environment; and
- user safety.

Article 4.3.9.

Types of aquaculture establishments and equipment

Aquaculture establishments and equipment differ widely in their characteristics. This section presents some considerations for effective disinfection of different types of aquaculture establishments and equipment.

1. Ponds

   Ponds are generally large and may be earthen based or be fitted with plastic liners. These characteristics together with the large volumes of water make cleaning prior to decontamination difficult and high organic loads may affect many chemical disinfectants. Ponds should be drained of water and have as much organic matter as possible removed prior to disinfection. All water and organic matter should be disinfected or disposed of in a biosecure manner. Earthen ponds should be dried thoroughly and lime compounds applied to raise pH and aid the inactivation of pathogenic agents. Scraping, ploughing or tilling of the base of unlined ponds will also aid in incorporation of liming compounds and drying.

2. Tanks

   Tank construction material (e.g. fibreglass, concrete or plastic) will determine the type of disinfection method used. Bare concrete tanks are susceptible to corrosion by acids and potential damage by high pressure sprayers. They are also porous and therefore require longer application of chemicals to ensure disinfection. Plastic, painted and fibreglass tanks are more easily disinfected because they have smooth, non-porous surfaces that facilitate thorough cleaning and are resistant to most chemicals.

   Tanks should be drained of water and have as much organic matter as possible removed prior to disinfection. Water and organic matter should be disinfected or disposed of in a biosecure manner. Tank equipment should be removed for separate cleaning and disinfection, and all organic waste and debris removed. Tank surfaces should be washed using high-pressure sprayers or mechanical scrubbing with detergent to remove fouling such as algae and biofilms. Heated water may be used to enhance the cleaning process. Before application of disinfectants any excess cleaning water should be drained and disinfected or disposed of in a biosecure manner.

   When disinfectants are applied to vertical surfaces, care should be taken to ensure that adequate contact time is maintained before the disinfectant is drained. Following disinfection, tanks should be rinsed to remove all residues and allowed to dry completely.
3. **Pipes**

*Disinfection* of pipes may be difficult due to lack of access. Pipe construction material should be taken into consideration when selecting the *disinfection* method.

Pipes can be cleaned through the use of alkaline or acid solutions, or foam projectile pipe cleaning systems. For cleaning to be effective, biofilms must be removed followed by flushing of the resulting particulate matter and thorough rinsing.

Once pipes are cleaned, chemical *disinfectants* or circulation of heated water can be used. All steps require pipes to be fully filled so that internal surfaces are treated.

4. **Cage nets and other fibrous materials**

Nets used in cage culture are often large, difficult to handle, have significant levels of biofouling and are usually made from fibrous materials that trap organic matter and moisture. Nets should be dedicated to a single *aquaculture establishment* or area because they have a high likelihood of contamination and may be difficult to disinfect.

Once the net has been removed from the water, it should be transferred directly to the net washing site. Nets should be thoroughly cleaned prior to *disinfection* to remove organic matter and aid in the penetration of chemical *disinfectants*. Cleaning of nets is best achieved by first removing gross biofouling and then washing with a detergent solution. Water and organic matter should be disposed of in a biosecure manner.

Following cleaning, nets may be disinfected by complete immersion in chemical *disinfectants* or heated water. Treatment duration should be sufficient to allow penetration into net material. Treatment may have a detrimental impact upon the strength of nets. This must be considered when deciding upon the treatment method to be applied to ensure net integrity is not compromised. Following *disinfection*, nets should be dried before storage. If rolled nets are not completely dry they will retain moisture which may enhance survival of the *pathogenic agent*.

Other fibrous materials such as wood, ropes and dip nets have characteristics similar to cage nets and they require special consideration. Wherever possible, it is recommended that equipment is site specific if it includes fibrous material.

5. **Vehicles**

The likelihood of *vehicle* contamination will be determined by their use, e.g. transportation of mortalities, live *aquatic animals*, harvested *aquatic animals*. All potentially contaminated internal and external surfaces should be disinfected. Special consideration should be given to areas likely to be contaminated such as the internal surface of *containers*, pipes, transportation water and waste. The application of corrosive *disinfectants* to *vehicles* should be avoided or if used, corrosive residues removed following *disinfection* by thorough rinsing. Oxidative compounds such as chlorines are the most commonly used *disinfectants* for *vehicles*.

All boats should undergo routine *disinfection* to ensure that they do not transfer *pathogenic agents*. The level of contamination of boats will be determined by their use. Boats used to harvest or to remove dead *aquatic animals* from *aquaculture* sites should be considered as highly likely to be contaminated. Organic material should be regularly removed from decks and work areas.

As part of the *disinfection* planning process, an assessment should be made to identify areas likely to be contaminated such as in and around machinery, holding tanks, bilges and pipes. All loose equipment should be removed, cleaned and disinfected separately from *disinfection* of the boat. Additional procedures should be developed for well-boats because of their potential to transfer *pathogenic agents* through the discharge of contaminated water. Contaminated effluent water should be disinfected prior to discharge (refer to Article 4.3.11.).

Where possible, boats should be placed on land or dry-docked, for *disinfection* in order to limit waste water entering the aquatic environment and to allow access to hull and niche areas. Biofouling organisms, that may act as *vectors*, and fomites should be removed.

Where boats cannot be removed to land or dry-docked, a *disinfection* method should be chosen that minimises the discharge of toxic chemicals into the aquatic environment. Divers should inspect and clean hulls. Where appropriate, mechanical methods such as high-pressure sprayers or steam cleaners should be considered as an alternative to chemical *disinfection* for cleaning above and below the water-line. Fumigation may also be considered for large areas if they can be adequately sealed.
Chapter 4.3.- Disinfection of aquaculture establishments and equipment

6. Buildings

Aquaculture establishments include buildings for culture, harvesting and processing of aquatic animals, and other buildings associated with storage of feed and equipment.

The approach to disinfection may vary depending on the structure of the building and degree of contact with contaminated material and equipment.

Buildings should be designed to allow effective cleaning and thorough application of disinfectants to all internal surfaces. Some buildings will contain complex piping, machinery and tank systems that may be difficult to disinfect. Wherever possible, buildings should be cleared of debris and emptied of equipment, prior to disinfection.

Misting or foaming agents are options for disinfection of complex areas and vertical surfaces. Fumigation can be considered for large or difficult to access areas if buildings can be adequately sealed.

7. Containers

Containers range from simple plastic bins used to transport harvested aquatic animal products or dead aquatic animals through to complex tank systems used for the transport of live aquatic animals.

Containers are generally manufactured using smooth non-porous material (e.g. plastic, stainless steel) which can be easily disinfected. They should be considered high risk items because they are in close contact with aquatic animals or their products (e.g. blood, diseased aquatic animals). In addition the need to move them between locations makes them potential fomites for the spread of pathogenic agents. In the case of transport of live aquatic animals, containers may also have pipes and pumping systems and confined spaces that should also be disinfected.

All water should be drained from the container and any aquatic animals, faecal matter and other organic material removed by flushing with clean water and disposed of in a biosecure manner. All pipes and associated pumps should also be inspected and flushed. Containers should then be washed using appropriate chemical detergents combined with high-pressure water cleaners or mechanical scrubbing.

All internal and external surfaces of containers should be treated using an appropriate disinfection method. They should then be rinsed and inspected to ensure there are no organic residues and stored in a manner that allows them to drain and dry quickly.

8. Biofilters

Biofilters associated with closed or semi-closed production systems are an important control point for disease. Biofilters are designed to maintain a colony of beneficial bacteria to enhance water quality. The conditions that support these bacteria may also enhance survival of some pathogenic agents should they be present. It is normally not possible to disinfect biofilters without also destroying beneficial bacteria. Therefore potential water quality issues should be taken into account when planning strategies for disinfection of biofilters.

When disinfecting biofilters and their substrates, the system should be drained, organic residues removed and surfaces cleaned. Disinfection of biofilter systems can be undertaken by modifying water pH levels (using either acid or alkaline solutions). Where this is undertaken, the pH levels must be sufficient to inactivate the pathogenic agent, but should not be corrosive to pumps and equipment within the biofilter system. Alternatively, the biofilter can be completely dismantled, including removal of biofilter substrate, and the components cleaned and disinfectants applied separately. In the case of emergency disease response, the latter procedure is recommended. The biofilter substrate should be replaced if it cannot be effectively disinfested. Biofilter systems should be thoroughly rinsed before re-stocking.

9. Husbandry and harvesting equipment

Aquaculture establishments will normally have a range of husbandry and harvesting equipment that come into close contact with aquatic animals and have potential to act as fomites. Examples include graders, automatic vaccinators and fish pumps.

The general principles described in Article 4.3.4. should be applied to disinfection of husbandry and harvesting equipment. Each item should be examined to identify areas that come into close contact with aquatic animals and where organic material accumulates. If required, equipment should be dismantled to allow adequate cleaning and application of disinfectants.
Article 4.3.10.

Personal equipment

Disinfection of personal equipment should consider the likelihood and degree of contamination associated with previous use. Where possible, personal equipment should be site specific to avoid the need for regular disinfection.

Equipment should be chosen which is non-absorbent and easy to clean. All staff entering a production area should use protective clothing that is clean and uncontaminated. On entry and exit of production areas boots should be cleaned and disinfected. When footbaths are used they should incorporate a cleaning procedure to remove accumulations of organic material and mud, be sufficiently deep to cover boots, use a disinfectant solution that is not inactivated by organic matter and be regularly refreshed with a new solution.

Some types of personal equipment such as dive equipment may require special attention because they are difficult to disinfect, may be moved from site to site and are often prone to chemical corrosion. Frequent rinsing of equipment will assist in reducing build-up of organic matter and make disinfection more efficient. Equipment should be allowed to dry thoroughly to ensure that moist microenvironments that may harbour pathogenic agents are minimised.

Article 4.3.11.

Disinfection of water

Aquaculture establishments may need to disinfect intake and effluent water to eliminate pathogenic agents. The most appropriate disinfection method will differ depending on the disinfection objective and the characteristics of the water to be disinfected.

Exclusion of aquatic animals and removal of suspended solids from the water to be treated are essential prior to the application of disinfectants. Pathogens are known to adhere to organic and inorganic matter and removal of suspended solids can significantly reduce loading of pathogenic agents in water. Removal of suspended solids can be achieved by filtration or settlement of suspended material. The most suitable filtration system will depend on the initial quality of water, volumes to be filtered, capital and operating costs and reliability.

Physical (e.g. UV irradiation) and chemical (e.g. ozone, chlorine and chlorine dioxide) disinfectants are commonly used to disinfect water. Suspended solids should be removed prior to the application of these disinfectants because organic matter may inhibit oxidative disinfection processes and suspended solids inhibit UV transmittance, reducing the efficacy of UV irradiation. A combination of methods may be beneficial where they are synergistic or where a level of redundancy is required.

It is essential to monitor the efficacy of water disinfection. This can be achieved by direct testing for pathogenic agents of concern, indirect monitoring of indicator organisms or monitoring of residual levels of disinfectants.

Management of chemical residues is important to avoid toxic effects on aquatic animals. For example, residuals formed between ozone and seawater such as bromide compounds are toxic to early life stages of aquatic animals and may be removed using charcoal filtration. Residual chlorine should be removed from water by chemical deactivation or off gassing.

NB: FIRST ADOPTED IN 2009; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 4.4.

RECOMMENDATIONS FOR SURFACE DISINFECTION OF SALMONID EGGS

Article 4.4.1.

Introduction

The practice of disinfecting salmonid eggs at hatcheries is an essential part of ensuring that pathogenic agents are not transferred between incubators and between facilities and forms a part of routine hatchery hygiene protocols. The disinfection process is also important for international trade in salmonid eggs between countries, zones or compartments to prevent the transfer of some pathogenic agents. Although generally effective for disinfection of the egg surface and reproductive fluids, the use of disinfectants will not prevent vertical transmission.

Salmonid eggs may be disinfected with a number of chemical agents. However, the most common method used is disinfection with the iodine-based product, povidone-iodine.

Iodophores, commonly povidone-iodine solutions, have the advantage of providing a neutral pH, being non-irritant and are relatively non-toxic. The neutral pH is important for minimising toxicity and ensuring efficacy. It is recommended to follow manufacturer’s instructions to identify circumstances where pH may be a concern. If other iodine based agents are used for disinfection it is essential that they be adequately buffered.

Article 4.4.2.

Disinfection protocol for salmonid eggs

This disinfection protocol may be applied to newly fertilised or eyed salmonid eggs. However newly fertilised eggs should be allowed to commence hardening prior to undergoing the disinfection protocol. Although there is a considerable margin of safety for hardened eggs, the disinfection protocol is not recommended for unfertilised ova or during fertilisation. It is essential that the pH of the iodophore solution is maintained between 6 and 8.

To disinfect salmonid eggs the following protocol should be applied:

1) rinse in pathogen-free 0.9% to 1.1% saline (30-60 seconds) to remove organic matter; then

2) immerse in an iodophore solution containing 100 ppm available iodine for a minimum of 10 minutes. The iodophore concentration should be monitored to ensure effective levels are maintained. The ratio of eggs to iodophore solution should be a maximum of 1:4; then

3) rinse again in pathogen-free 0.9% to 1.1% saline for 30-60 seconds; then

4) hold in pathogen-free water.

All rinsing and disinfection solutions should be prepared using pathogen free water. Iodophore solutions may be buffered using sodium bicarbonate (NaHCO₃) if the pH is low.

NB: FIRST ADOPTED IN 2015; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 4.5.

CONTINGENCY PLANNING

Article 4.5.1.

A number of diseases are regarded as posing a potential threat to aquaculture as well as to wild stocks of aquatic animals world-wide. The introduction of such diseases into countries recognised to be free from these diseases or into countries with an established control system and eradication programme for such diseases, may result in significant losses. In order to diminish such losses, the Competent Authority responsible for aquatic animal health may need to act quickly and should develop a contingency plan(s) before such events occur.

Article 4.5.2.

Legal powers

Countries must establish the necessary legal provisions that are needed for the implementation of a contingency plan(s). Such legal powers must include provisions for establishing a list of diseases for which action is needed, definitions of how such diseases should be managed if detected, provisions for access to infected/suspected sites, and other legal provisions, as needed.

Article 4.5.3.

Crises centre(s)

Countries must establish specified crises centre(s) (disease control centre[s]) that shall have the responsibility for the co-ordination of all control measures to be carried out. Such centres could either be located centrally or locally, depending on the infrastructure in a given country. A list of the crises centre(s) that has(have) the necessary facilities to carry out disease control measures should be made widely available.

The contingency plan(s) should also state that the crises centre(s) has(have) the authority to act rapidly to bring a given disease situation under control by contacting the personnel, organisations, aquaculture establishments, etc., that are involved directly or indirectly in managing an outbreak of a disease.

Article 4.5.4.

Personnel

The contingency plan(s) should provide information on the staff required to undertake the control measures, their responsibilities, and instructions on the chain of command.

Article 4.5.5.

Instructions

Countries establishing a contingency plan(s) should provide a detailed set of instructions on actions to be taken when a specified aquatic animal disease is suspected or confirmed. These could include:

1) diagnostic procedures in national reference laboratories;
2) confirmation of diagnosis, if necessary, at an OIE Reference Laboratory;
3) standing instructions to aquatic animal health personnel in the field;
4) instructions for handling/disposal of dead aquatic animals at an aquaculture establishment;
5) instructions for sanitary slaughtering;
Chapter 4.5. - Contingency planning

6) instructions for disease control at the local level;
7) instructions for the establishment of quarantine areas and observation (surveillance) zones;
8) provisions for controlling movements of aquatic animals in established zones;
9) disinfection procedures;
10) falling procedures;
11) surveillance methods for establishing successful eradication;
12) re-stocking procedures;
13) compensation issues;
14) reporting procedures;
15) provisions for raising public awareness of aquatic animal disease.

Article 4.5.6.

Diagnostic laboratories

Countries establishing a contingency plan(s) should establish national reference laboratories having the necessary facilities for diagnostic work on aquatic animal diseases that can be carried out rapidly. The national laboratory(ies) must also have established a set of instructions as regards rapid transportation of samples, and established protocols for quality assurance and diagnostic procedures to be used.

Article 4.5.7.

Training programmes

Countries establishing a contingency plan(s) must establish necessary training programmes to ensure that skills in field, administrative and diagnostic procedures are maintained. Announced and unannounced field exercises for administrators and aquatic animal personnel should be carried out to maintain the state of readiness.

NB: FIRST ADOPTED IN 2000.
CHAPTER 4.6.

FALLOWING IN AQUACULTURE

Article 4.6.1.

Introduction

Gaps in aquaculture production at the same location are commonly recognised to be of value in resting or restoring the local environment. As part of this strategy, fallowing can break re-infection cycles by removing loci of a disease from a farm. Consequently, fallowing is often carried out as a regular disease management measure in aquaculture, especially prior to the introduction of new populations of aquatic animals into a previously used site. In order to promote improved health in aquaculture, the Aquatic Animal Health Service in a country may encourage the use of fallowing as a routine management strategy for many diseases. Account should be taken of the likely beneficial effects of fallowing in proportion to the economic costs involved. The Aquatic Animal Health Service should also consider such factors as the level of risk to the local and national aquaculture operations, previous knowledge of the severity of a disease(s), the infective period and distribution of the pathogenic agent(s), the socioeconomic conditions, and benefits pertaining to the general aquatic resources. When the infective period is not known, the farm may be fallowed for a period, the length of which should be based on a risk assessment.

However, where an official stamping-out policy is being carried out for a disease of concern, the Aquatic Animal Health Service should require that an infected aquaculture establishment, and all other aquaculture establishments in an officially established infected zone, be subjected to a required period of fallowing, if necessary synchronised.

Article 4.6.2.

Legal powers

In cases where fallowing may be a compulsory measure, for instance in the establishment or restoration of a disease free zone, countries should establish a legal framework for the implementation of fallowing procedures in aquaculture establishments. Legal provisions could include:

1) defining the disease circumstances when fallowing or synchronised fallowing is required;

2) defining mechanisms based on risk assessment where individual disease-specific measures may be determined, including disinfection and the length of the fallowing period prior to the re-introduction of susceptible species;

3) following permission by the Competent Authority to restock with susceptible species, defining a period of surveillance and diagnostic to verify freedom from the specified disease.

Article 4.6.3.

Technical parameters for the implementation of a statutory fallowing plan

Fallowing of a farm should start immediately after:

1) removal of all susceptible species of aquatic animals for the disease of concern; and
2) removal of all species capable of acting as vectors of the disease of concern; and
3) if appropriate, removal of other species; and
4) removal of water in which infected stocks have been held, where feasible; and
5) equipment and other materials contaminated or otherwise capable of harbouring infection have either been removed or subjected to disinfection to standards approved by the Aquatic Animal Health Service.

The length of the statutory fallowing period should be based on scientific evidence of the likelihood of a pathogenic agent remaining infective outside its aquaculture host(s) in the local environment, at a level likely to cause an unacceptable risk of re-infection of the aquaculture establishment. Account should be taken of the extent of the disease outbreak, local availability of alternative hosts, the survival and infectivity characteristics of the pathogenic agent and the local climatological, geographical and hydrographical factors. In addition, the level of risk to the local aquaculture industry and
wider aquatic resources may be included. A scientifically based risk assessment approach should be used to determine the length of the fallowing period.

Article 4.6.4.

Instructions

Countries establishing fallowing procedures should develop a detailed set of instructions for disinfection of aquaculture establishments prior to fallowing. For this purpose, the instructions set out in Chapter 4.3. of the Aquatic Code and in Chapter 1.1.3. of the Aquatic Manual should be used as guidelines, taking into account current scientific knowledge on the efficacy of the treatments for the pathogenic agent of concern.

Article 4.6.5.

Restocking

No aquaculture establishment that has been under compulsory fallowing should be restocked until the fallowing period has been completed and permission from the Competent Authority has been received. When restocking, care should be taken not to use stocks of aquatic animals that would compromise the objectives of the fallowing procedure.

To increase confidence in the effectiveness of the fallowing procedures, all farms subjected to compulsory fallowing should have a period of high level official surveillance after susceptible species have been restocked. The duration and intensity of the surveillance should be appropriate for the disease of concern and local conditions.

NB: FIRST ADOPTED IN 2003; MOST RECENT UPDATE ADOPTED IN 2016.
CHAPTER 4.7.

HANDLING, DISPOSAL AND TREATMENT OF AQUATIC ANIMAL WASTE

Article 4.7.1.

Introduction

The objective of this chapter is to provide guidance on storage, transport, disposal and treatment of aquatic animal wastes so as to manage risks to aquatic animal health. The recommendations in this chapter are general in nature. The choice of one or more of the recommended methods should comply with relevant local and national legislation.

Disposal methods should take into consideration a range of factors, including the cause of mortality. It may be appropriate to carry out a risk assessment on the disposal options.

In the case of killing of animals for disease control purposes or unusually large mortalities, this may require approval from, or supervision by, the Competent Authority.

In the event of aquatic animal mortalities of a significant nature in aquaculture or in the wild, the Competent Authority should be notified so that necessary steps can be taken to dispose of the dead aquatic animals, in order to minimise the risk for possible spread of aquatic animal disease.

Article 4.7.2.

Scope

The scope of this chapter covers aquatic animal waste derived from: i) routine aquaculture operations; ii) on shore processing, irrespective of origin; iii) mass killing for disease control purposes and iv) mass mortality (including in the wild).

Article 4.7.3.

Definitions

Aquatic animal waste means the entire body or parts of aquatic animals that have died or been killed for disease control purposes as well as slaughtered aquatic animals, and their parts, that are not intended for human consumption.

High risk waste means aquatic animal waste that constitutes, or is suspected of constituting, a serious health risk to aquatic animals or humans.

Low risk waste means aquatic animal waste that is not high risk waste.

Article 4.7.4.

Governance

The Competent Authority should oversee the efficient and effective disposal of aquatic animal waste. Cooperation among all relevant agencies and stakeholders involved in aquatic animal health is necessary to ensure safe handling and disposal. In this context the following aspects should be addressed:

1) physical, logistical and data access by relevant personnel, in cooperation with stakeholders, including access of the Competent Authority to the aquatic animal waste;

2) movement controls and the authority to make exemptions under certain biosecurity conditions, for example for transport of aquatic animal waste to another location for disposal;
3) the determination of the method and location of disposal, and the necessary equipment and facilities, by the Competent Authority, in consultation with other authorities including government organisations responsible for the protection of human health and the environment.

Article 4.7.5.

Storage, transport and labelling

Following collection, aquatic animal waste should be stored for the minimum time practical; however, where storage is necessary there should be sufficient capacity for the expected waste and the Competent Authority may require additional measures.

The storage area should be separated from aquaculture sites and bodies of water to minimise the risk of spread of pathogenic agents. The containers of stored aquatic animal waste should be leak-proof and secured to prevent contact with aquatic animals, other animals or birds and unauthorised personnel.

Aquatic animal waste infected by an agent causing a disease referred to in the Aquatic Code or suspected of being so, may not be transported without permission from the Competent Authority. The Competent Authority may assess the requirement for this condition based on the disease situation in the Member Country (e.g. where a disease referred to in the Aquatic Code is enzootic in the Member Country).

If low risk waste becomes contaminated with high risk waste, such waste should then be considered high risk waste.

Containers used for transport of aquatic animal waste should be leak-proof and labelled regarding content. Transport should be accompanied by appropriate documentation detailing origin, content and destination to allow tracing if required.

Equipment used for transportation should be cleaned and disinfected before being returned, as described in Chapter 4.3.

Article 4.7.6.

Approval and operational requirements of disposal plants

1. Requirement for approval
   All disposal plants dealing with aquatic animal waste should be approved by the Competent Authority. However, disposal plants using only low risk waste for production of products not intended to be used in animals may be exempted from approval but should be registered by the Competent Authority.

2. Conditions for approval
   For a disposal plant to be approved to deal with aquatic animal waste, it should:
   a) be adequately separated from thoroughfares through which contamination may be spread, other premises (such aquaculture facilities, slaughterhouses, processing plants) and bodies of water, so as to minimise the risk of spread of pathogenic agents;
   b) be designed and equipped to the satisfaction of the Competent Authority;
   c) have access to approved or accredited laboratories;
   d) fulfil requirements for handling the aquatic animal waste and products specified by the Competent Authority.
   Any substantial proposed changes to the disposal plant should be approved by the Competent Authority.
   Approval should be withdrawn or suspended, as appropriate, if a disposal plant no longer fulfils the criteria given by the Competent Authority.

3. Operating requirements
   The disposal plant should operate using procedures that minimise the risk of spread of pathogenic agents, including:
   a) separation of clean and unclean areas, including consideration of workflow, and good hygienic procedures for personnel;
   b) equipment and surfaces should be easy to clean and disinfect;
   c) handling and treatment of aquatic animal waste should take place as soon as possible after being received;
   d) effluent waste water should be collected and disinfected before leaving the premises;
Chapter 4.7.- Handling, disposal and treatment of aquatic animal waste

e) incorporating measures to prevent access of birds, insects, rodents or other animals to the disposal plant;
f) a system for registration and labelling of material for tracing purposes.

A system for internal control, identifying critical points and means of control for such points, should be in place at the disposal plants. A general documentation system for internal control including sampling for control of critical points should be established.

Spot checks of batches should be carried out to check the microbiological standards following processing. Products from incineration plants may be exempted from such checks. The Competent Authority may grant exemptions on specified conditions.

If testing of the product from processed high risk waste shows that the product is not satisfactorily produced and thus poses a risk for the spread of pathogenic agents, disposal plants should report immediately to the Competent Authority who may then require additional measures. These products should not be transported from disposal plants without permission from the Competent Authority.

Results from the different samples and checks should be kept for a given period decided upon by the Competent Authority. Analyses and sampling should be carried out in accordance with international standards.

Disposal plants applying treatments based on time and pressure should be able to measure and record these parameters.

Disposal plants should maintain records related to quantity and type of raw material received, supplier, quantity and type of finished product, receivers, critical check points, and deviations from provisions stipulated in relevant regulations. These should be made available to the Competent Authority on request.

Article 4.7.7.

Methods for disposal of high risk waste

Recommended methods for disposal of high risk waste from aquatic animals are as follows:

1. Rendering

Rendering will inactivate all of the known aquatic animal pathogenic agents.

Rendering is generally carried out in a closed system using a combination of mechanical treatments and time/temperature combinations leading to stable, sterilised products, such as fish meal and fish oil.

The process typically involves pre-heating to 50–60°C, followed by cooking of the raw waste at 95–100°C for 15 to 20 minutes. The oil and proteins are separated by pressing and centrifuging involving temperatures of 90°C. The production of meal involves further high temperature treatments.

2. Incineration

Incineration is a controlled burning process carried out in fixed incinerators or mobile air curtain incinerators. Mobile air curtain incinerators enable the process to be carried out on site thus removing the need to transport the aquatic animal waste.

Incinerators may only be capable of handling limited volumes of aquatic animal waste.

3. Sterilisation

The minimum requirement for sterilisation is a core temperature of at least 90°C for at least 60 minutes, but other time/temperature combinations are also available and effective.

4. Composting

Composting does not inactivate all pathogenic agents; therefore, high risk waste should be heated (85°C for 25 minutes or an equivalent temperature/time combination) prior to the composting process.

Effective composting depends upon a combination of pH, temperature, moisture and time factors. Depending on the type of composting (e.g. windrows, closed vessel) and the raw material used, as well as the climatic conditions, the temperature parameters of the process and the heat distribution in the material may be different.

When held in windrows, the entire material needs an exposure time of at least two weeks at 55°C, while in closed vessels exposure to 65°C for one week is required.
5. **Biogas production**

Biogas production does not inactivate all *pathogenic agents*; therefore, high risk waste should be treated to ensure inactivation of *pathogenic agents* prior to the biogas production process. The method chosen should be shown to inactivate the *pathogenic agents* of concern.

Biogas production is a process whereby organic matter in biological waste products is fermented under anaerobic conditions.

The two main types of biogas production are mesophilic anaerobe digestion and thermophilic anaerobe digestion. Both processes are normally continuous, and a portion of the end material is removed every 2–12 hours. There is a risk that new material which has been in the reactor for only 2–12 hours may be removed with the finished products.

6. **Ensiling**

Ensiling does not inactivate all *pathogenic agents*; therefore, high risk waste should be heated (85°C for 25 minutes or an equivalent temperature/time combination) prior to the ensiling process.

Ensiling of *aquatic animal* waste in an organic acid such as formic acid is an effective method of inactivating most *pathogenic agents* within 48 hours. The pH in the ensiling process should be maintained at, or below, 4.0 throughout the process.

7. **Burial**

Burial may take place either in a landfill site or other locations approved by the Competent Authority based on risk assessments as regards aquatic animal health, public health and possible environmental impacts.

Whenever possible, the *aquatic animal* waste should be subjected to a treatment that ensures inactivation of the *pathogenic agents* prior to burial.

In selecting an acceptable burial site, consideration should be given to the following:

- **a)** Location – the possible effects of the fire’s heat, smoke and odour on nearby structures, underground and aerial utilities, roads and residential areas. The site should be surrounded by an adequate firebreak.
- **b)** Access – easy access for equipment and delivery of aquatic animal waste. Fencing and restricted admittance may be necessary.
- **c)** Pit construction – rocky areas should be avoided. Soils with good stability, capable of withstanding the weight of equipment used to dig and fill the pits, should be selected. If required, diversion banks can be constructed to prevent surface runoff entering the pit or to prevent any liquids escaping from the burial site. Pit dimensions depend on the volume of the *aquatic animal* waste to be buried and should be easy to fill.
- **d)** Pit closure – contents should be covered with unslaked lime (CaO) at a rate of 85 kg per 1,000 kg of aquatic animal waste to hasten decomposition and prevent scavenging.

8. **Pyre-burning**

Pyre-burning may not be suitable for large amounts of *aquatic animal* waste.

In selecting an acceptable pyre-burning site, the following considerations are important:

- **a)** Location – the possible effects of the fire’s heat, smoke and odour on nearby structures, underground and aerial utilities, roads and residential areas. The site should be surrounded by an adequate firebreak.
- **b)** Access – for equipment to construct the pyre and maintain the fire, for the delivery of fuel and *aquatic animal* waste.

Pyre-burning needs considerable amounts of fuel and all required fuel should be on site before the burning is started. If the pyre-burning is carried out correctly, *aquatic animal* wastes will be destroyed within 48 hours.

When leaving the pyre-burning site, vehicles and containers should be disinfected.

Alternatively, high risk waste may be disposed of by any methods, approved by the Competent Authority, which ensure an equivalent reduction of risk.

**Article 4.7.8.**

**Methods of disposal for low risk waste**

Low risk waste can be disposed of using all methods described in Article 4.7.7. In the case of composting or biogas production it is not necessary to heat treat the low risk waste prior to disposal.
Alternatively, the following methods may be used:

1. **Ensiling**
   Ensiling of aquatic animal waste in an organic acid such as formic acid is an effective method of inactivating most pathogenic agents within 48 hours. The pH in the ensiling process should be maintained at, or below, 4.0 throughout the process.
   
   The Competent Authority may require ensiling as a treatment prior to one of the disposal methods described in Article 4.7.7.

2. **Pasteurisation**
   Pasteurisation does not inactivate all pathogenic agents. Heat treatment at temperatures below 100°C can be considered as pasteurisation. Pasteurisation may use a range of time/temperature combinations.

In addition, the Competent Authority may permit low risk waste to be disposed of by other means, or used for any other purposes, following an assessment of the risk from such methods or uses.

**Article 4.7.9.**

**Mass mortality events**

Mass mortality of aquatic animals can arise from natural events or killing for disease control purposes (refer to Chapter 7.4.). This may lead to the need for disposal of large numbers of dead aquatic animals and is often subject to intense public and media scrutiny. The Competent Authority should conduct disposal operations within acceptable scientific principles that will address the risks of spread of the pathogenic agent, and public and environmental concerns.

1. **Preparedness**
   Successful disposal with minimum delay is achieved by advance planning and preparation:
   
   a) Preparedness planning should engage other relevant government agencies and stakeholders such as industry organisations, animal welfare organisations, emergency response organisations, and media.
   
   b) Standard operating procedures should be developed (including documented decision-making processes, training of staff).
   
   c) Pre-arranged mechanisms to access emergency funding for the disposal operation.
   
   d) Information sharing with officials involved in the disposal operation, stakeholders, politicians and the media is essential. A well informed spokesperson should be available at all times to answer enquiries.
   
   e) Resource readiness planning should address such items as personnel, transport, storage facilities, equipment, fuel, protective clothing and logistical support. Special equipment, such as well boats, may be required.

2. **Critical elements**
   Critical elements which need to be considered in planning and implementation include:
   
   a) rapid disposal of the dead aquatic animals;
   
   b) methods of treatment and disposal should address capacity issues and the risks of spread of pathogenic agents;
   
   c) adequate funding and staff resources;
   
   d) addressing the risk of spread of pathogenic agents by vectors and fomites;
   
   e) stakeholder cooperation;
   
   f) safety of personnel;
   
   g) environmental concerns;
   
   h) societal acceptance.
Chapter 4.7.- Handling, disposal and treatment of aquatic animal waste

3. Choice of disposal methods

The Competent Authority may determine the dead aquatic animals to be either high risk waste or low risk waste and select an appropriate disposal method in accordance with the risk (refer to Articles 4.7.7. and 4.7.8.). Should the chosen disposal option be applied near the border of a neighbouring country, the Competent Authority of that country should be informed.

NB: FIRST ADOPTED IN 2010.
CHAPTER 4.8.

CONTROL OF PATHOGENIC AGENTS IN AQUATIC ANIMAL FEED

Article 4.8.1.

Introduction

Feed can be a source of infectious disease in aquatic animals.

Because aquatic animals are often a principal ingredient in feed for aquatic animals, and because the use of semi-processed, raw and live feed continues to be a common practice, the risk of disease transmission via feed should be addressed.

Article 4.8.2.

Purpose and scope

The purpose of this chapter is to address transmission of infectious diseases of aquatic animals via feed to prevent entry and spread into a country, zone or compartment free from pathogenic agents of concern.

This chapter applies to the production and use of all products destined for feed and feed ingredients whether produced commercially or on farm.

Risk analysis principles (in accordance with Chapter 2.1.) should be applied to determine the risks associated with the production and use of feed in aquatic animals.

This chapter is complementary to guidance provided by the Codex Code of Practice on Good Animal Feeding (CAC/RCP 54-2004).

Article 4.8.3.

Responsibilities

The responsibilities of the Competent Authority include setting and enforcing regulatory requirements related to animal feed, and verifying that these requirements are met. This also includes raising awareness about risks related to use of unprocessed or semi-processed feed in aquaculture.

Feed producers have the responsibility to ensure that production of feed is performed in a manner to prevent the spread of diseases of aquatic animals. Records and contingency plans should be in place, as appropriate, to enable the tracing, recall, or destruction of non-compliant products. All personnel involved in the harvest, manufacture, transport, storage and handling of feed and feed ingredients should be adequately trained and aware of their role and responsibility in preventing the spread of infectious diseases of aquatic animals. Equipment for producing, storing and transporting feed and feed ingredients should be kept clean and maintained in good working order.

Owners and managers of aquaculture establishments should adhere to regulatory requirements and implement biosecurity plans on their farms in order to manage risks related to the use of semi-processed, raw and live feed. This can be done through identification of disease free sources and record keeping for traceability purposes, implementation of on farm risk mitigation measures, and early detection of infectious diseases.

Private veterinarians and other aquatic animal health professionals providing specialist services to producers and to the feed industry may be required to meet specific regulatory requirements pertaining to the services they provide (e.g. disease reporting, quality standards, transparency).
Chapter 4.8.- Control of pathogenic agents in aquatic animal feed

Article 4.8.4.

Hazards associated with aquatic animal feed

Biological hazards that may be present in feed and feed ingredients include pathogenic agents such as bacteria, viruses, fungi, and parasites. The scope of these recommendations covers listed diseases and other pathogenic agents that cause an adverse effect on aquatic animal health.

Chemical and physical hazards associated with feed and feed ingredients are not addressed in this chapter.

Antimicrobial resistance arising from the use of antimicrobial agents in feed is addressed in Section 6.

Article 4.8.5.

Risk pathways and exposure

Feed may be contaminated with pathogenic agents present at the time of harvesting, transport, storage and processing of commodities used as feed ingredients. Contamination may also occur during manufacture, transport, storage and use of feed. Poor hygienic practices during processing and manufacture, transport and storage are potential sources of contamination with pathogenic agents.

Aquatic animals can be directly exposed to pathogenic agents in feed. Aquatic animals can also be indirectly exposed through contamination of the environment by feed.

Article 4.8.6.

Risk management

1. Use of safe feed and feed ingredients

Some commodities undergo significant processing such as heat treatment, acidification, extrusion and extraction. There may be a negligible likelihood that pathogenic agents will survive in such products if they have been produced in accordance with Good Manufacturing Practice.

Criteria provided in Chapter 5.4. may be used to assess the safety of commodities to be used as feed or feed ingredients.

Articles X.X.3. of all disease-specific chapters in Sections 8 to 11 list commodities considered safe for any purpose including use as feed or feed ingredients.

Competent Authorities should also consider sourcing feed and feed ingredients from a country, zone or compartment free from pathogenic agents of concern.

2. Use of feed and feed ingredients from sources that may not be free from pathogenic agents of concern

When using feed and feed ingredients from sources that may not be free from pathogenic agents of concern, Competent Authorities should consider the following risk mitigation measures:

a) treatment (e.g. by heating or acidification) of the commodity using a method approved by the Competent Authority to inactivate pathogenic agent(s) as per Articles X.X.10. (for Chapter 10.4. the relevant Article is 10.4.14.) of all disease-specific chapters in Sections 8 to 11; or

b) confirmation (e.g. by testing) that pathogenic agents are not present in the commodity; or

c) use of feed only in populations that are not susceptible to the pathogenic agent(s) in question and where susceptible species will not come into contact with the feed or its waste products.

3. Feed production

To prevent contamination by pathogenic agents during processing, manufacture, storage and transport of feed and feed ingredients, the following is recommended:

a) flushing, sequencing or physical cleaning-out of manufacturing lines and storage facilities should be performed between batches as appropriate;

b) buildings and equipment for processing and transporting feed and feed ingredients should be constructed in a manner that facilitates hygienic operation, maintenance and cleaning and prevents contamination;
c) *feed* manufacturing plants should be designed and operated in a manner that avoids cross-contamination between batches;

d) processed *feed* and *feed ingredients* should be stored separately from unprocessed *feed ingredients*, under appropriate storage conditions;

e) *feed* and *feed ingredients*, manufacturing equipment, storage facilities and their immediate surroundings should be kept clean;

f) measures to inactivate *pathogenic agents*, such as heat treatment, should be used where appropriate;

g) labelling should provide for the identification of *feed* and *feed ingredients* as to the batch, place and date of production to assist in tracing *feed* and *feed ingredients*.

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NB: FIRST ADOPTED IN 2008; MOST RECENT UPDATE ADOPTED IN 2015.
SECTION 5.
TRADE MEASURES, IMPORTATION/EXPORTATION PROCEDURES AND HEALTH CERTIFICATION

CHAPTER 5.1.
GENERAL OBLIGATIONS RELATED TO CERTIFICATION

Article 5.1.1.

A combination of factors should be taken into account to facilitate international trade in aquatic animals and aquatic animal products, without incurring unacceptable risks to human and aquatic animal health.

Because of differences between countries in their aquatic animal health situations, various options are offered by the Aquatic Code. The aquatic animal health situation in the exporting country, in the transit country or countries and in the importing country should be considered before determining the requirements for trade. To maximise harmonisation of the aquatic animal health aspects of international trade, Competent Authorities of Member Countries should base their import requirements on OIE standards. These requirements should be included in the certificates drawn up in accordance with the model international aquatic animal health certificates provided for in Chapter 5.11.

Certificates should be exact and concise, and should clearly address the requirements of the importing country. For this purpose, prior consultation between Competent Authorities of importing and exporting countries may be necessary. This consultation helps to determine the exact requirements of the certification.

Certificates should be issued and signed by a certifying official authorised by the Competent Authority to perform inspections, and endorsed through signature and/or official stamp of the Competent Authority. The certification requirements should not include conditions for diseases that are not transmitted by the commodity concerned. The certificate should be signed in accordance with the provisions of Chapter 5.2.

When officials of a Competent Authority wish to visit another country for matters of professional interest to the Competent Authority of the other country, the latter should be informed prior to any such visit. This visit should be mutually agreed upon between Competent Authorities.

Article 5.1.2.

Responsibilities of the importing country

1) The import requirements included in the international aquatic animal health certificate should assure that commodities introduced into the importing country comply with OIE standards. Importing countries should align their requirements with the recommendations in the relevant standards of the OIE. If there are no such recommendations or if the country chooses a level of protection requiring measures more stringent than the standards of the OIE, these should be based on an import risk analysis conducted in accordance with Chapter 2.1.
Chapter 5.1.- General obligations related to certification

2) The international aquatic animal health certificate should not include requirements for the exclusion of pathogenic agents or aquatic animal diseases that are present in the importing country and are not subject to any official control programme. The measures imposed on imports to manage the risks posed by a pathogenic agent or aquatic animal disease should not be more stringent than those applied as part of the official control programme operating within the importing country.

3) The international aquatic animal health certificate should not include measures against pathogenic agents or diseases that are not OIE listed, unless the importing country has demonstrated through an import risk analysis, carried out in accordance with Section 2, that the pathogenic agent or disease poses a significant risk to the importing country.

4) The transmission of the requirements of the importing country or certificates from the Competent Authority of the importing country and the communication of import requirements to persons other than the Competent Authority of another country necessitates that copies of these documents be also sent to the Competent Authority of the exporting country. This important procedure avoids delays and difficulties that may arise between traders and Competent Authorities when the authenticity of the certificates or permits is not established.

The transmission of this information is the responsibility of Competent Authorities of the exporting country. However, it can be issued by private sector veterinarians at the place of origin of the commodities when this practice is the subject of appropriate approval and authentication by Competent Authorities.

5) Situations may arise that result in changes to the consignee, identification of the means of transportation, or frontier post after a certificate is issued. If it is determined that these do not change the aquatic animal health or public health status of the consignment, then they should not prevent the acceptance of the certificate.

Article 5.1.3.

Responsibilities of the exporting country

1) An exporting country should, on request, supply the following to importing countries:
   a) information on the aquatic animal health situation and national aquatic animal health information systems to determine whether that country is free or has zones or compartments free from listed diseases, and on the pathway followed to achieve disease freedom e.g. historical freedom, absence of susceptible species or targeted surveillance, including the regulations and procedures in force to maintain the free status;
   b) regular and prompt information on the occurrence of listed diseases;
   c) details of the country's ability to apply measures to control and prevent listed diseases;
   d) information on the structure of the Competent Authority and the authority that they exercise;
   e) technical information, particularly on biological tests and vaccines applied in all or part of the country.

2) Competent Authorities of exporting countries should:
   a) have official procedures for the authorisation of certifying officials, defining their functions and duties as well as conditions of oversight and accountability, including possible suspension and termination of the authorisation;
   b) ensure that relevant instructions and training are provided to certifying officials;
   c) monitor the activities of the certifying officials to verify their integrity and impartiality.

3) The Competent Authority of the exporting country is ultimately accountable for certification used in international trade.

Article 5.1.4.

Responsibilities in case of an incident related to importation

1) International trade involves a continuing ethical responsibility. Therefore, if within a reasonable period subsequent to an export taking place, the Competent Authority becomes aware of the appearance or reappearance of a disease that has been specifically included in the international aquatic animal health certificate or other disease of potential epidemiological importance to the importing country there is an obligation for the Competent Authority to notify the importing country, so that the imported commodities may be inspected or tested and appropriate action be taken to limit the spread of the disease should it have been inadvertently introduced.
2) If a disease appears in aquatic animals and is associated with importation of commodities, the Competent Authority of the exporting country should be informed. This will enable the exporting country to investigate as this may be the first available information on the occurrence of the disease in a previously free aquatic animal population. The Competent Authority of the importing country should be informed of the result of the investigation because further action may be required if the source of the infection did not originate in the exporting country.

3) In case of suspicion, on reasonable grounds, that an international aquatic animal health certificate may be fraudulent, the Competent Authorities of the importing country and exporting country should conduct an investigation. Consideration should also be given to notifying any third country that may have been implicated. All associated consignments should be kept under official control, pending the outcome of the investigation. Competent Authorities of all countries involved should fully cooperate with the investigation. If the international aquatic animal health certificate is found to be fraudulent, every effort should be made to identify those responsible so that appropriate action can be taken in accordance with the relevant legislation.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 5.2.

CERTIFICATION PROCEDURES

Article 5.2.1.

Protection of the professional integrity of the certifying official

Certification should be based on the highest possible ethical standards, the most important of which is that the professional integrity of the certifying official should be respected and safeguarded.

It is essential to include in the certificate only those specific statements that can be accurately and honestly signed by a certifying official. For example, these requirements should not include certification of an area as being free from diseases that are not notifiable in that country, or the occurrence of which the signing certifying official is not necessarily informed about. It is unacceptable to ask for certification for events that will take place after the document is signed when these events are not under the direct control and supervision of the signing certifying official.

Article 5.2.2.

Certifying officials

Certifying officials should:

1) be authorised by the Competent Authority of the exporting country to sign international aquatic animal health certificates;
2) only certify matters that are within their own knowledge at the time of signing the certificate, or that have been separately attested by another party authorised by the Competent Authority;
3) sign only at the appropriate time certificates that have been completed fully and correctly; where a certificate is signed on the basis of supporting documentation, the certifying official should have verified or be in possession of that documentation before signing;
4) have no conflict of interest in the commercial aspects of the aquatic animals or aquatic animal products being certified and be independent from the commercial parties.

Article 5.2.3.

Preparation of international aquatic animal health certificates

Certificates should be drawn up in accordance with the following principles:

1) Certificates should be designed so as to minimise the potential for fraud including use of a unique identification number, or other appropriate means to ensure security. Paper certificates should bear the signature of the certifying official and the official identifier (stamp) of the issuing Competent Authority. Each page of a multiple page certificate should bear the unique certificate number and a number indicating the number of the page out of the total number of pages. Electronic certification procedures should include equivalent safeguards.
2) Certificates should be written using terms that are simple, unambiguous and as easy to understand as possible, without losing their legal meaning.
3) If so required, certificates should be written in the language of the importing country. In such circumstances, they should also be written in a language understood by the certifying official.
4) Certificates should require appropriate identification of aquatic animals and aquatic animal products except where this is impractical (e.g. eyed eggs).
5) Certificates should not require a certifying official to certify matters that are outside his/her knowledge or that he/she cannot ascertain and verify.
6) Where appropriate, when presented to the certifying official, certificates should be accompanied by notes of guidance indicating the extent of enquiries, tests or examinations expected to be carried out before the certificate is signed.
Chapter 5.2. - Certification procedures

7) The text of a certificate should not be amended except by deletions that should be signed and stamped by the certifying official.

8) The signature and stamp should be in a colour different to that of the printing of the certificate. The stamp may be embossed instead of being a different colour.

9) Only original certificates should be accepted by the importing country.

10) Replacement certificates may be issued by a Competent Authority to replace original certificates that have been, for example, lost, damaged, contain errors, or where the original information is no longer correct. These replacements should be provided by the issuing authority and be clearly marked to indicate that they are replacing the original certificate. A replacement certificate should reference the number and the issue date of the certificate that it supersedes. The superseded certificate should be cancelled and where possible, returned to the issuing authority.

Article 5.2.4.

Electronic certification

1) Certification may be provided by electronic exchange of data sent directly from the Competent Authority of the exporting country to the Competent Authority of the importing country.
   a) Systems providing electronic certificates normally provide an interface with the commercial organisation marketing the commodity for provision of information to the certifying authority. The certifying official should have access to all necessary information such as origin of aquatic animals and laboratory results.
   b) When exchanging electronic certificates and in order to fully utilise electronic data exchange, Competent Authorities should use internationally standardised language, message structure and exchange protocols. Guidance for electronic certification in standardised Extensible Markup Language (XML) as well as secure exchange mechanisms between Competent Authorities is provided by the United Nations Centre for Trade Facilitation and Electronic Business (UN/CEFACT).
   c) A secure method of electronic data exchange should be ensured by digital authentication of the certificates, encryption, non-repudiation mechanisms, controlled and audited access and firewalls.

2) Electronic certificates should carry the same information as conventional certificates.

3) The Competent Authority should have in place systems for the security of electronic certificates against access by unauthorised persons or organisations.

4) The certifying official should be officially responsible for the secure use of his/her electronic signature.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2015.
CHAPTER 5.3.

OIE PROCEDURES RELEVANT TO THE AGREEMENT ON THE APPLICATION OF SANITARY AND PHYTOSANITARY MEASURES OF THE WORLD TRADE ORGANIZATION

Article 5.3.1.

The Agreement on the Application of Sanitary and Phytosanitary Measures and role and responsibility of the OIE

The Agreement on the Application of Sanitary and Phytosanitary Measures (SPS Agreement) encourages the Members of the World Trade Organization to base their sanitary measures on international standards, guidelines and recommendations, where they exist. Members may choose to adopt a higher level of protection than that provided by international texts if there is a scientific justification or if the level of protection provided by the relevant international texts is considered to be inappropriate. In such circumstances, Members are subject to obligations relating to risk assessment and to a consistent approach of risk management.

The SPS Agreement encourages Governments to make a wider use of risk analysis: WTO Members shall undertake an assessment as appropriate to the circumstances of the actual risk involved.

The SPS Agreement, in Article 7, obliges WTO Members to notify changes in, and provide relevant information on, sanitary measures which may, directly or indirectly, affect international trade.

The SPS Agreement recognises the OIE as the relevant international organisation responsible for the development and promotion of international animal health standards, guidelines, and recommendations affecting trade in live animals and animal products, including aquatic animals and their products.

Article 5.3.2.

Introduction on the judgement of the equivalence of sanitary measures

The importation of aquatic animals and aquatic animal products involves a degree of risk to the aquatic animal health status of an importing country. The estimation of that risk and the choice of the appropriate risk management option(s) are made more difficult by differences among the aquatic animal health and production systems in Member Countries. It is now recognised that significantly different aquatic animal health and production systems can provide equivalent aquatic animal and human health protection for the purpose of international trade, with benefits to both the importing country and the exporting country.

These recommendations are to assist Member Countries to determine whether sanitary measures arising from different aquatic animal health and production systems may provide the same level of aquatic animal and human health protection. They discuss principles which might be utilised in a judgement of equivalence, and outline a step-wise process for trading partners to follow in facilitating a judgement of equivalence. These provisions are applicable whether equivalence applies at the level of specific measures or on a systems-wide basis, and whether equivalence applies to specific areas of trade or commodities, or generally.

Article 5.3.3.

General considerations on the judgement of the equivalence of sanitary measures

Before trade in aquatic animals or their products may occur, an importing country must be satisfied that its aquatic animal health status will be appropriately protected. In most cases, the risk management measures drawn up will rely in part on judgements made about the aquatic animal health and production system(s) in the exporting country and the effectiveness of sanitary procedures undertaken there. Systems operating in the exporting country may differ from those in the importing country and from those in other countries with which the importing country has traded. Differences may
be with respect to infrastructure, policies and/or operating procedures, laboratory systems, approaches to the pests and diseases present, border security and internal movement controls.

International recognition of the legitimacy of different approaches to achieving the importing country's appropriate level of protection (ALOP) has led to the principle of equivalence being included in trade agreements, including the SPS Agreement of the WTO.

Benefits of applying equivalence may include:
1) minimising costs associated with international trade by tailoring aquatic animal health measures to local circumstances;
2) maximising aquatic animal health outcomes for a given level of resource input;
3) facilitating trade by achieving the required health protection through less trade restrictive sanitary measures; and
4) decreased reliance on relatively costly commodity testing and isolation procedures in bilateral or multilateral agreements.

The Aquatic Code recognises equivalence by recommending alternative sanitary measures for many diseases and pathogenic agents. Equivalence may be gained, for example, by enhanced surveillance and monitoring, by the use of alternative test, treatment or isolation procedures, or by combinations of the above. To facilitate the judgement of equivalence, Member Countries should base their sanitary measures on the standards, guidelines and recommendations of the OIE.

It is essential to apply a scientific risk analysis to the extent practicable in establishing the basis for a judgement of equivalence.

Article 5.3.4.

Prerequisite considerations in a judgement of equivalence

1. Application of risk assessment

Application of the discipline of risk assessment provides a structured basis for judging equivalence among different sanitary measures as it allows a close examination to be made of the effect of a measure(s) on a particular step(s) in the importation pathway, and the relative effects of proposed alternative measure(s) on the same or related steps.

A judgement of equivalence needs to assess the sanitary measure in terms of its effectiveness regarding the particular risk or group of risks against which the measure is designed to protect. Such an assessment may include the following elements: the purpose of the measure, the level of protection achieved by the measure and the contribution the measure makes to achieving the ALOP of the importing country.

2. Categorisation of sanitary measures

Proposals for equivalence may be in terms of a measure comprising a single component of a measure (e.g. an isolation procedure, a test or treatment requirement, a certification procedure) or multiple components (e.g. a production system for commodity), or a combination of measures. Multiple components or combinations of measures may be applied consecutively or concurrently.

Sanitary measures are those described in each chapter of the Aquatic Code which are used for risk reduction and are appropriate for particular diseases. Sanitary measures may be applied either alone or in combination and include test requirements, processing requirements, inspection or certification procedures, quarantine confinements, and sampling procedures.

For the purposes of judging equivalence, sanitary measures can be broadly categorised as:

a) infrastructure: including the legislative base (e.g. aquatic animal health law) and administrative systems (e.g. organisation of national and regional aquatic animal health authorities, emergency response organisations);

b) programme design/implementation: including documentation of systems, performance and decision criteria, laboratory capability, and provisions for certification, audit and enforcement;
Chapter 5.3.- OIE procedures relevant to the Agreement on the Application of Sanitary and Phytosanitary Measures of the World Trade Organization

5.3.5. Principles for judgement of equivalence

In conjunction with the above considerations, judgement of the equivalence of sanitary measures should be based on application of the following principles:

1) an importing country has the right to set the level of protection it deems appropriate (its ALOP) in relation to human and animal life and health in its territory; this ALOP may be expressed in qualitative or quantitative terms;

2) the importing country should be able to describe the reason for each sanitary measure i.e. the level of protection intended to be achieved by application of the identified measure against a hazard;

3) an importing country should recognise that sanitary measures different from the ones it has proposed may be capable of providing the same level of protection;

4) the importing country should, upon request, enter into consultations with the exporting country with the aim of facilitating a judgement of equivalence;

5) any sanitary measure or combination of sanitary measures can be proposed for judgement of equivalence;

6) an interactive process should be followed that applies a defined sequence of steps, and utilises an agreed process for exchange of information, so as to limit data collection to that which is necessary, minimise administrative burden, and facilitate resolution of claims;

7) the exporting country should be able to demonstrate objectively how the alternative sanitary measure(s) proposed as equivalent will provide the same level of protection;

8) the exporting country should present a submission for equivalence in a form that facilitates judgement by the importing country;

9) the importing country should evaluate submissions for equivalence in a timely, consistent, transparent and objective manner and in accordance with appropriate risk assessment principles;

10) the importing country should take into account any knowledge of and prior experience with the Veterinary Authority or other Competent Authority of the exporting country;

11) the exporting country should provide access to enable the procedures or systems which are the subject of the equivalence judgement to be examined and evaluated upon request of the importing country;

12) the importing country should be the sole determinant of equivalence, but should provide to the exporting country a full explanation for its judgement;

13) to facilitate a judgement of equivalence, Member Countries should base their sanitary measures on relevant OIE standards;

14) to allow the judgement of equivalence to be reassessed if necessary, the importing country and the exporting country should keep each other informed of significant changes to infrastructure, health status or programmes which may bear on the judgement of equivalence; and

15) an importing country should give positive consideration to a request by an exporting developing country for appropriate technical assistance that would facilitate the successful completion of a judgement of equivalence.
Article 5.3.6.

Sequence of steps to be taken in judgement of equivalence

There is no single sequence of steps which must be followed in all judgements of equivalence. The steps that trading partners choose will generally depend on the circumstances and their trading experience. The interactive sequence of steps described below may be useful for all sanitary measures irrespective of their categorisation as infrastructure, programme design/implementation or specific technical requirement components of an aquatic animal health and production system.

This sequence assumes that the importing country is meeting its obligations under the WTO SPS Agreement and has in place a transparent measure based either on an international standard or a risk analysis.

Recommended steps are:

1) the exporting country identifies the measure(s) for which it wishes to propose an alternative measure(s), and requests from the importing country a reason for its sanitary measure in terms of the level of protection intended to be achieved against a hazard(s);

2) the importing country explains the reason for the measure(s), in terms which would facilitate comparison with an alternative sanitary measure(s) and consistent with the principles set out in these provisions;

3) the exporting country demonstrates the case for equivalence of an alternative sanitary measure(s) in a form which facilitates analysis by an importing country;

4) the exporting country responds to any technical concerns raised by the importing country by providing relevant further information;

5) judgement of equivalence by the importing country takes into account as appropriate:
   a) the impact of biological variability and uncertainty;
   b) the expected effect of the alternative sanitary measure(s) on all relevant hazards;
   c) OIE standards;
   d) application of solely qualitative frameworks where it is not possible or reasonable to conduct quantitative risk assessment;

6) the importing country notifies the exporting country of its judgement and the underlying reasons within a reasonable period of time:
   a) recognition of the equivalence of the exporting country's alternative sanitary measure(s);
   b) request for further information; or
   c) rejection of the case for equivalence of the alternative sanitary measure(s);

7) an attempt should be made to resolve any differences of opinion over judgement of a case, either interim or final, by using an agreed mechanism to reach consensus (e.g. the OIE informal procedure for dispute mediation), or by referral to an agreed expert;

8) depending on the category of measures involved, the importing country and the exporting country may enter into a formal equivalence agreement giving effect to the judgement or a less formal acknowledgement of the equivalence of a specific measure(s) may suffice.

An importing country recognising the equivalence of an exporting country's alternative sanitary measure(s) needs to ensure that it acts consistently with regard to applications from third countries for recognition of equivalence applying to the same or very similar measure(s). Consistent action does not mean however that a specific measure(s) proposed by several exporting countries should always be judged as equivalent as a measure(s) should not be considered in isolation but as part of a system of infrastructure, policies and procedures.

Article 5.3.7.

Sequence of steps to be taken in establishing a zone/compartment and having it recognised for international trade purposes

There is no single sequence of steps which should be followed in establishing a zone or a compartment. The steps that the Veterinary Services or Aquatic Animal Health Services of the importing country and the exporting country choose and implement will generally depend on the circumstances existing within the countries and at their borders, and their trading history. The recommended steps are:
Chapter 5.3.- OIE procedures relevant to the Agreement on the Application of Sanitary and Phytosanitary Measures of the World Trade Organization

1. For zoning
   a) The exporting country identifies a geographical area within its territory, which it considers to contain an aquatic animal subpopulation with a distinct health status with respect to a specific disease/specific diseases, based on surveillance.  
   b) The exporting country describes in the biosecurity plan for the zone the measures which are being, or will be, applied to distinguish such an area epidemiologically from other parts of its territory, in accordance with the recommendations in the Aquatic Code.
   c) The exporting country provides:
      i) the above information to the importing country, with an explanation of why the area can be treated as an epidemiologically separate zone for international trade purposes;
      ii) access to enable the procedures or systems that establish the zone to be examined and evaluated upon request by the importing country.
   d) The importing country determines whether it accepts such an area as a zone for the importation of aquatic animals and aquatic animal products, taking into account:
      i) an evaluation of the exporting country’s Veterinary Services or Aquatic Animal Health Services;
      ii) the result of a risk assessment based on the information provided by the exporting country and its own research;
      iii) its own aquatic animal health situation with respect to the disease(s) concerned; and
      iv) other relevant OIE standards.
   e) The importing country notifies the exporting country of its determination and the underlying reasons, within a reasonable period of time, being:
      i) recognition of the zone; or
      ii) request for further information; or
      iii) rejection of the area as a zone for international trade purposes.
   f) An attempt should be made to resolve any differences over recognition of the zone, either in the interim or finally, by using an agreed mechanism to reach consensus such as the OIE informal procedure for dispute mediation (Article 5.3.8.).
   g) The Veterinary Authorities or other Competent Authorities of the importing and exporting countries should enter into a formal agreement recognising the zone.

2. For compartmentalisation
   a) Based on discussions with the relevant industry, the exporting country identifies within its territory a compartment comprising an aquatic animal subpopulation contained in one or more establishments or other premises operating under common management practices related to biosecurity. The compartment contains an identifiable aquatic animal subpopulation with a distinct health status with respect to specific disease(s). The exporting country describes how this status is maintained through a partnership between the relevant industry and the Veterinary Authority or other Competent Authority of the exporting country.
   b) The exporting country examines the compartment's biosecurity plan and confirms through an audit that:
      i) the compartment is epidemiologically closed throughout its routine operating procedures as a result of effective implementation of its biosecurity plan; and
      ii) the surveillance and monitoring programme in place is appropriate to verify the status of such a subpopulation with respect to such disease(s).
   c) The exporting country describes the compartment, in accordance with the recommendations in the Aquatic Code.
   d) The exporting country provides:
      i) the above information to the importing country, with an explanation of why such a subpopulation can be treated as an epidemiologically separate compartment for international trade purposes; and
      ii) access to enable the procedures or systems that establish the compartment to be examined and evaluated upon request by the importing country.
   e) The importing country determines whether it accepts such a subpopulation as a compartment for the importation of aquatic animals and aquatic animal products, taking into account:
      i) an evaluation of the exporting country's Veterinary Services or Aquatic Animal Health Services;
      ii) the result of a risk assessment based on the information provided by the exporting country and its own research;
iii) its own aquatic animal health situation with respect to the disease(s) concerned; and
iv) other relevant OIE standards.

f) The importing country notifies the exporting country of its determination and the underlying reasons, within a reasonable period of time, being:
i) recognition of the compartment; or
ii) request for further information; or
iii) rejection of such a subpopulation as a compartment for international trade purposes.

g) An attempt should be made to resolve any differences over recognition of the compartment, either in the interim or finally, by using an agreed mechanism to reach consensus such as the OIE informal procedure for dispute mediation (Article 5.3.8.).

h) The Veterinary Authorities or other Competent Authorities of the importing and exporting countries should enter into a formal agreement recognising the compartment.

i) The Veterinary Authority or other Competent Authority of the exporting country should promptly inform importing countries of any occurrence of a disease in respect of which the compartment was defined.

Article 5.3.8.

The OIE informal procedure for dispute mediation

OIE shall maintain its existing voluntary in-house mechanisms for assisting Member Countries to resolve differences. In-house procedures which will apply are that:

1) Both parties agree to give the OIE a mandate to assist them in resolving their differences.

2) If considered appropriate, the Director General of the OIE recommends an expert, or experts, and a chairman, as requested, agreed by both parties.

3) Both parties agree on the terms of reference and working programme, and to meet all expenses incurred by the OIE.

4) The expert or experts are entitled to seek clarification of any of the information and data provided by either country in the assessment or consultation processes, or to request additional information or data from either country.

5) The expert or experts shall submit a confidential report to the Director General of the OIE, who will transmit it to both parties.

NB: FIRST ADOPTED IN 2013.
CHAPTER 5.4.

CRITERIA TO ASSESS THE SAFETY OF AQUATIC ANIMAL COMMODITIES

In the context of this chapter the word ‘safety’ is applied only to animal health considerations for listed diseases.

Article 5.4.1.

Criteria to assess the safety of aquatic animals and aquatic animal products for any purpose from a country, zone or compartment not declared free from disease X

In all disease chapters, point 1 of Article X.X.3. lists aquatic animals and aquatic animal products that can be traded for any purpose from a country, zone or compartment not declared free from disease X. The criteria for inclusion of aquatic animals and aquatic animal products in point 1 of Article X.X.3. are based on the absence of the pathogenic agent in the traded aquatic animals and aquatic animal products or inactivation of the pathogenic agent by treatment or processing.

The assessment of the safety of the aquatic animals and aquatic animal products using the criteria relating to treatment or processing can only be undertaken where treatments or processing are well defined. It may not be necessary to provide details of the entire treatment or process undertaken. However, the steps considered critical in the inactivation of the pathogenic agent of concern should be detailed.

It is assumed that treatment or processing (i) uses standardised protocols, which include the steps considered critical in the inactivation of the pathogenic agent of concern; (ii) is conducted in accordance with Good Manufacturing Practices; and (iii) that any other steps in the treatment, processing and subsequent handling of the aquatic animal product do not jeopardise the safety of the traded aquatic animal product.

Criteria

For an aquatic animal or aquatic animal product to be considered safe for international trade under the provisions of Article X.X.3., it should comply with the following criteria:

1) Absence of pathogenic agent in the traded aquatic animal or aquatic animal product:
   a) There is strong evidence that the pathogenic agent is not present in the tissues from which the aquatic animal or aquatic animal product is derived.

   AND

   b) The water (including ice) used to process or transport the aquatic animal or aquatic animal product is not contaminated with the pathogenic agent and the processing prevents cross contamination of the aquatic animal or aquatic animal product to be traded.

OR

2) Even if the pathogenic agent is present in, or contaminates the tissues from which the aquatic animal or aquatic animal product is derived, the treatment or processing to produce the aquatic animal or aquatic animal product to be traded inactivates the pathogenic agent:
   a) physical (e.g. temperature, drying, smoking);

   AND/OR

   b) chemical (e.g. iodine, pH, salt, smoke);

   AND/OR

   c) biological (e.g. fermentation).
Article 5.4.2.

Criteria to assess the safety of aquatic animals or aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from disease X

In all disease chapters, point 1 of Article X.X.12. (amphibian and fish disease chapters) and Article X.X.11. (crustacean and mollusc disease chapters) lists aquatic animals or aquatic animal products for retail trade for human consumption. The criteria for inclusion of aquatic animals or aquatic animal products in point 1 of Article X.X.12. (amphibian and fish disease chapters) and Article X.X.11. (crustacean and mollusc disease chapters) include consideration of the form and presentation of the product, the expected volume of waste tissues generated by the consumer and the likely presence of viable pathogenic agent in the waste.

For the purpose of this criterion retail means the selling or provision of aquatic animals or aquatic animal products directly to the consumer with the intended purpose of human consumption. The retail pathway may also include wholesale distribution of the products provided they are not further processed by the wholesale distributor or the retailer, i.e. are not subjected to actions such as gutting, cleaning, filleting, freezing, thawing, cooking, unpacking, packing or repackaging.

It is assumed that: (i) the aquatic animals or aquatic animal products are used for human consumption only; (ii) waste may not always be handled in an appropriate manner that mitigates the introduction of the pathogenic agent; the level of risk is related to the waste disposal practices in each Member’s country or territory; (iii) treatment or processing prior to importation is conducted in accordance with Good Manufacturing Practices, and (iv) any other steps in the treatment, processing and subsequent handling of the aquatic animals or aquatic animal products prior to importation do not jeopardise the safety of the traded aquatic animals or aquatic animal products.

Criteria

For aquatic animals or aquatic animal products to be considered for international trade under the provisions of point 1 of Article X.X.12. (amphibian and fish disease chapters) and Article X.X.11. (crustacean and mollusc disease chapters), it should comply with the following criteria:

1) The aquatic animal or aquatic animal product is prepared and packaged for retail trade for human consumption; AND

EITHER

2) it includes only a small amount of raw waste tissues generated by the consumer;

OR

3) the pathogenic agent is not normally found in the waste tissues generated by the consumer.

NB: FIRST ADOPTED IN 2009; MOST RECENT UPDATE ADOPTED IN 2011.
CHAPTER 5.5.

CONTROL OF AQUATIC ANIMAL HEALTH RISKS ASSOCIATED WITH TRANSPORT OF AQUATIC ANIMALS

Article 5.5.1.

General considerations

1) These considerations should be used as recommendations when countries introduce measures to control the aquatic animal health risks related to the transport of these aquatic animals and aquatic animal products. These recommendations do not address aquatic animal welfare.

2) Vehicles (or containers) used for the transport of aquatic animals shall be designed, constructed and fitted in such a way as to withstand the weight of the aquatic animals and water and to ensure their safety during transportation. Vehicles shall be thoroughly cleansed and disinfected before use in accordance with the recommendations given in the Aquatic Code.

3) Vehicles (or containers) in which aquatic animals are confined during transport shall be secured to maintain optimal conditions for the aquatic animals during transport, and to allow easy access by the attendant.

Article 5.5.2.

Particular considerations for containers

1) The construction of containers intended for transportation of aquatic animals shall be such that the accidental release of water, etc., is prevented during transport.

2) In the case of the transportation of aquatic animals, provision shall be made to enable preliminary observation of the contents of containers.

3) Containers in transit in which there are aquatic animal products shall not be opened unless the Aquatic Animal Health Service of the transit country consider it necessary. If this is the case, containers shall be subject to precautions to prevent contamination.

4) Containers shall be loaded only with one kind of product or, at least, with products not susceptible to contamination by one another.

5) It rests with each country to decide on the facilities it requires for the transport and importation of aquatic animals and aquatic animal products in containers.

Article 5.5.3.

Particular considerations for the transport of aquatic animals by air

1) The stocking densities for the transport of aquatic animals in containers should be determined by taking the following into consideration when transporting by air:
   a) the total volume of available space for each type of aquatic animal;
   b) the oxygenation capacity available to supply the containers while on the ground and during all stages of the flight.

   With regard to fish, molluscs and crustaceans, the space reserved for each aquatic animal species in containers that have been fitted for the separate transportation of several aquatic animals or for the transportation of groups of aquatic animals should comply with acceptable densities specified for the species in question.

2) The OIE approved International Air Transport Association (IATA) Regulations for live animals may be adopted if they do not conflict with national legislative arrangements. (Copies of these Regulations are obtainable from IATA, 800 Place Victoria, P.O. Box 113, Montreal, Quebec H4Z 1M1, Canada.)
Chapter 5.5.- Control of aquatic animal health risks associated with transport of aquatic animals

Article 5.5.4.

**Disinfection and other sanitary measures**

1) *Disinfection* and all zoo-sanitary work should be carried out in order to:

   a) avoid all unjustified inconvenience and to prevent damage or injury to the health of people and aquatic animals;

   b) avoid damage to the structure of the vehicle or its appliances;

   c) prevent, as far as possible, any damage to aquatic animal products.

2) On request, the Aquatic Animal Health Service shall issue the transporters with a certificate indicating the measures that have been applied to all vehicles, the parts of the vehicle that have been treated, the methods used and the reasons that led to the application of the measures.

   In the case of aircraft, the certificate may be replaced, on request, by an entry in the General Declaration of the aircraft.

3) Likewise, the Aquatic Animal Health Service shall issue on request:

   a) a certificate showing the date of arrival and departure of the aquatic animals;

   b) a certificate to the shipper or exporter, the consignee and transporter or their representatives, indicating the measures applied.

Article 5.5.5.

**Treatment of transportation water**

Water to be used for transportation of aquatic animals should be appropriately treated after transport and/or before discharge in order to minimise the risk of transferring pathogens. The specific recommendations are provided in the chapter of the Aquatic Code on disinfection.

During transportation of aquatic animals, the transporter should not be permitted to evacuate and replace the water in the transport tanks except on specifically designated sites in the national territory. The waste and rinsing water should not be emptied into a drainage system that is directly connected to an aquatic environment where aquatic animals are present. The water from the tanks should therefore either be disinfected by a recognised process (for example, 50 mg iodine or chlorine/litre for one hour), or sprayed over land that does not directly drain into waters containing aquatic animals. Each country shall designate the sites in their national territories where these operations can be carried out.

Article 5.5.6.

**Discharge of infected material**

The Aquatic Animal Health Service shall take all practical measures to prevent the discharge of any untreated infective material, including transport water, into internal or territorial waters.

Article 5.5.7.

**Particular considerations for the transport of live fish by well boat**

A well boat is a boat with integrated tanks to carry live fish in sea water that may operate with open valves to allow exchange of sea water. Therefore, well boats can present a biosecurity risk if the fish being carried are infected. Well boats are inherently difficult to disinfect.

1) Only healthy fish showing no clinical signs of disease on the day of loading should be transported. The well boat must have the capability of fully closed containment of fish during its operation if so required.

2) The stocking densities should be determined by taking both the total volume of available space for each species of fish and the oxygenation/aeration capacity available to supply the fish during all stages of transport into consideration.

3) Fish may be transported by well boat from an infected site if this is part of a disease response plan agreed to by the Competent Authority.

4) Provision shall be made to enable preliminary observation of the contents in the well, and monitoring equipment should be available where appropriate.
Chapter 5.5.- Control of aquatic animal health risks associated with transport of aquatic animals

5) Access by farm staff to the vessel and from the vessel to the farm cages, including the equipment, should be restricted.

6) Transporting fish of different health status at the same time increases the risk of disease transfer between those fish and is discouraged.

7) Well boats may exchange water in their tanks with the environment except in designated areas in proximity to aquaculture establishments or areas with protected wild populations. The Aquatic Animal Health Service should designate the areas based upon a risk assessment.

8) Multiple deliveries of fish during the same trip should be avoided. Where unavoidable the order of deliveries should be made to sites of a higher health status first (e.g. youngest year class), to a single aquaculture establishment, or establishments of the same health status.

9) In the event of mortality occurring during transport, a contingency plan capable of dealing with full containment and disposal of dead fish, via an approved disposal method, should be available. This plan should be prepared in accordance with the recommendations on handling and disposal of carcasses and wastes of aquatic animals (in preparation).

10) Well boats should not operate in adverse inclement weather conditions that may force the operation to divert from the planned route and schedule of transport.

11) The well boat should be cleaned and, where required, disinfected to an acceptable standard before re-use. The level of disinfection should be proportional to the risk. Well boats should maintain a disinfection checklist which should be kept with the ship’s log and should be open to audit. It is essential to ensure that all fish are removed from the system before cleaning. All organic matters should be removed through the process of cleaning before disinfection commences. The general principles and specific recommendations as outlined in the Aquatic Manual should be consulted for guidance.

12) When travelling between areas and zones of different health levels, cleaning and, if required, disinfection procedures should be followed and implemented to a standard approved by the Aquatic Animal Health Service.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2010.
CHAPTER 5.6.

AQUATIC ANIMAL HEALTH MEASURES
APPLICABLE BEFORE AND AT DEPARTURE

Article 5.6.1.

1) Each country should only authorise the exportation from its territory of live aquatic animals and aquatic animal products that are correctly identified, and inspected in accordance with the procedures outlined in the Aquatic Code and Aquatic Manual.

2) In certain cases, the above-mentioned aquatic animals could, in accordance with the wish of the importing country, be subjected to certain biological tests or to prophylactic parasitological procedures within limits of a defined period of time before their departure.

3) Observation of the above-mentioned aquatic animals before leaving the country may be carried out in the establishment where they were reared or at the frontier post. When they have been found to be clinically healthy and free from listed diseases or any other specified infectious disease by a member of the personnel of the Competent Authority or a certifying official approved by the importing country during the period of observation, the aquatic animals should be transported to the place of shipment in specially constructed containers, previously cleansed and disinfected, without delay and without coming into contact with other susceptible aquatic animals, unless these aquatic animals have health guarantees similar to those of the transported aquatic animals.

4) The transportation of aquatic animals for breeding or rearing or slaughter shall be carried out directly from the establishment of origin to the place of shipment or to the processing establishment in conformity with the conditions agreed between the importing and exporting countries.

Article 5.6.2.

Each country should only undertake the exportation of live aquatic animals or eggs or gametes destined for a country or zone or aquaculture establishment officially declared free from one or more of the listed diseases, when the exporting country or zone or aquaculture establishment of origin is itself officially declared free of the same disease(s). If the live aquatic animals originate in an infected aquaculture establishment or infected zone, with respect to the disease(s) in question, the exporting country should not export the aquatic animals if they have been exposed to infection by direct or indirect contact of a kind likely to cause transmission of the pathogenic agent(s), without the prior agreement of the importing country.

Article 5.6.3.

Each country exporting aquatic animals at any stage of development or aquatic animal products should inform the country of destination and when necessary the transit countries if, after exportation, diagnosis of a listed disease occurs in the establishment of origin, or in aquatic animals that were in the aquaculture establishment or natural water body at the same time as the exported animals, within a period of time that indicates that the exported consignment may have been infected.

Article 5.6.4.

Before the departure of the aquatic animals and aquatic animal products, a member of the personnel of the Competent Authority or a certifying official approved by the importing country should provide an international aquatic animal health certificate conforming with the models approved by the OIE (as shown in Chapter 5.11.) and worded in the languages agreed upon between the exporting country and the importing country and, when necessary, with the transit countries.
Article 5.6.5.

1) Before the departure of a consignment of aquatic animals on an international journey, the Competent Authority of the port, airport or district in which the frontier post is situated may, if it is considered necessary, have a health examination carried out on the consignment. The time and place of the examination shall be fixed taking into account customs and other formalities and in such a way as not to impede or unreasonably delay departure.

2) The Competent Authority referred to in point 1 above shall take necessary measures to:
   a) prevent the shipment of aquatic animals showing clinical signs of any listed disease;
   b) avoid entry into the container of possible vectors or pathogenic agents.

CHAPTER 5.7.

AQUATIC ANIMAL HEALTH MEASURES APPLICABLE DURING TRANSIT FROM THE PLACE OF DEPARTURE IN THE EXPORTING COUNTRY TO THE PLACE OF ARRIVAL IN THE IMPORTING COUNTRY

Article 5.7.1.

1) Any country through which the transit of aquatic animals has to be made, and that normally conducts commercial transactions with the exporting country, should not refuse the transit, subject to the reservations mentioned herein and on condition that notification is made of the proposed transit to the Competent Authority in charge of the frontier posts.

This notification shall state the species and quantities of aquatic animals, the methods of transport and the frontier posts of entry and exit in accordance with a previously arranged and authorised itinerary in the transit country.

2) Any country through which transit has to take place may refuse such transit if, in the exporting country or transit country that precedes it on the itinerary, certain diseases exist that have been specifically included in the international aquatic animal health certificates or in bilateral agreements. Alternatively, the Competent Authority of the transit country may impose conditions with regard to the method, including packaging, and route of transport.

3) Any transit country may require the presentation of international aquatic animal health certificates. Such a country may, in addition, cause an examination to be made by a member of the personnel of the Aquatic Animal Health Service on the health status of fish, molluscs or crustaceans in transit, except in cases where transport in sealed vehicles or containers is a condition of transit.

4) Any transit country may refuse passage through its territory of aquatic animals at one of its frontier posts if an examination carried out by a member of the personnel of the Aquatic Animal Health Service shows that the consignment of aquatic animals in transit is affected by or infected with any of the listed diseases and if these diseases are exotic to that country or the zone through which the transportation was to take place, or if there is an enforced control programme for the disease(s) in question, or if the international aquatic animal health certificate is inaccurate and/or unsigned or does not apply to fish, molluscs or crustaceans.

In these circumstances, the Competent Authority of the exporting country shall be informed immediately, thereby providing an opportunity for checking the findings or correcting the certificate.

If the diagnosis of any listed disease is confirmed or if the certificate cannot be corrected, the consignment of aquatic animals in transit shall either be returned to the exporting country if there is a common frontier with it, or be slaughtered or destroyed.

Article 5.7.2.

1) Any transit country may require vehicles used for the transit of aquatic animals through its territory to be constructed to prevent the escape and dispersion of waste water or other contaminated material.

2) Unloading of aquatic animals shall be permitted in the territory of the transit country only if an emergency situation arises. The importing country shall be informed of any unforeseen unloading in the transit country and the reason for it.

Article 5.7.3.

Vessels stopping in a port or passing through a canal or other navigable route situated in the territory of a country, on their way to a port situated in the territory of another country, must comply with the conditions required by the Competent Authority.
Chapter 5.7.- Aquatic animal health measures applicable during transit from the place of departure in the exporting country to the place of arrival in the importing country

Article 5.7.4.

1) If, for reasons beyond the control of its captain, a ship or aircraft calls or lands somewhere other than at a port or airport, or at a port or airport other than that at which it should normally call or land, the captain of the ship or aircraft, or his/her deputy, shall immediately notify the nearest Competent Authority or any other public authority of the new port of call or landing.

2) As soon as the Competent Authority is notified of this calling or landing place, it shall take appropriate action.

3) The aquatic animals on board the ship or aircraft shall not be permitted to leave the vicinity of the docking or landing place and the removal from the vicinity of any equipment or packing material accompanying them shall not be permitted.

4) When the measures prescribed by the Competent Authority have been carried out, the ship or aircraft shall be permitted, for aquatic animal health purposes, to proceed to the port or airport at which it would normally have called or landed or, if there are technical reasons for which this cannot be done, to a port or an airport that is more suitable.

NB: FIRST ADOPTED IN 1995.
CHAPTER 5.8.

FRONTIER POSTS IN THE IMPORTING COUNTRY

Article 5.8.1.

The Competent Authority shall provide specified frontier posts with an office comprising personnel, equipment and premises as the case may be and, in particular, means for:

1) detecting and isolating aquatic animal populations affected with or suspected of being affected with a disease;
2) carrying out disinfection of vehicles used to transport aquatic animals and aquatic animal products;
3) making clinical examinations and obtaining specimens of material for diagnostic purposes from live aquatic animals or carcasses of aquatic animals affected or suspected of being affected with a disease, and obtaining specimens of aquatic animal products suspected of contamination.

Furthermore, it is preferable that each port and international airport be provided with equipment for the sterilisation or incineration of any material dangerous to aquatic animal health.

Article 5.8.2.

When required by international traffic in transit, airports shall be provided, as soon as possible, with areas of direct transit; these must, however, comply with the conditions required by the Competent Authority.

Article 5.8.3.

Each Veterinary Authority shall keep at the disposal of the OIE Headquarters and any interested country on request:

1) a list of specified frontier posts and processing plants for aquatic animals in its territory that are approved for international trade;
2) the period of time required for notice to be given for the application of the arrangements contained in paragraph 2 of Articles 5.9.1. and 5.9.2.;
3) a list of airports in its territory that are provided with an area of direct transit.

CHAPTER 5.9.

AQUATIC ANIMAL HEALTH MEASURES APPLICABLE ON ARRIVAL

Article 5.9.1.

1) An importing country should only accept into its territory live aquatic animals that have been subjected to examination by a member of the personnel of the Aquatic Animal Health Service of the exporting country or a certifying official approved by the importing country and that are accompanied by an international aquatic animal health certificate (see Model Certificates given in Chapter 5.11.).

2) An importing country may require sufficient advance notification regarding the proposed date of entry into its territory of aquatic animals, stating the species, quantity, means of transport and the name of the frontier post. In addition, any importing country shall publish a list of the specified frontier posts supplied with the equipment required for conducting control operations at importation and enabling the importation and transit procedures to be carried out in the most speedy and efficacious way.

3) An importing country may prohibit the introduction into its territory of aquatic animals if these were found, on examination carried out at the frontier post by a member of the personnel of the Aquatic Animal Health Service, to be affected by a listed disease of concern to the importing country. Refusal of entry may also be applied to aquatic animals that are not accompanied by an international aquatic animal health certificate conforming to the requirements of the importing country. In these circumstances, the Competent Authority of the exporting country shall be informed immediately, thereby providing an opportunity for checking the findings or correcting the certificate.

   However, the importing country may prescribe that the importation be placed immediately in quarantine in order to carry out a clinical observation and biological examinations with a view to establishing a formal diagnosis. If the diagnosis of a listed disease is confirmed, or if the certificate cannot be corrected, the importing country may take the following measures:

   a) return the aquatic animals involved to the exporting country if this rejection does not involve transit through a third country;
   
   b) slaughter and destroy in cases where re-shipment would be dangerous from a health point of view or impossible from a practical point of view.

Article 5.9.2.

1) An importing country should only accept into its territory raw uneviscerated fish of those species susceptible to a listed disease destined for introduction into an aquatic environment or for human consumption that have been subjected to examination by a member of the personnel of the Aquatic Animal Health Service of the exporting country or a certifying official approved by the importing country, and that are accompanied by an international aquatic animal health certificate (see Model Certificates given in Chapter 5.11.).

2) An importing country may require sufficient advance notification regarding the proposed date of entry into its territory of a consignment of products of aquatic animal origin destined for human consumption, together with information on the nature, quantity and packaging of the products, as well as the name of the frontier post.

Article 5.9.3.

On arrival at a frontier post of a vehicle transporting aquatic animals infected with any specified listed disease, the vehicle shall be considered to be contaminated and the Aquatic Animal Health Service shall apply the following measures:

1) unloading of the vehicle and immediate transportation of any possibly contaminated material, such as water or ice, to an establishment assigned in advance for its destruction and the strict application of the aquatic animal health measures required by the importing country;
2) disinfection of:
   a) outer clothes and boots of the crew on the transporting vehicle;
   b) all parts of the vehicle that were used in the transport, moving and unloading of the aquatic animals.

CHAPTER 5.10.

MEASURES
CONCERNING INTERNATIONAL TRANSPORT OF
AQUATIC ANIMAL PATHOGENS AND
PATHOLOGICAL MATERIAL

Article 5.10.1.

Introduction

There is the risk that disease may occur as a result of the accidental release of aquatic animal pathogens during international transport of packaged materials. Such pathogens may already occur in the country or they may have been imported deliberately or inadvertently. It is therefore necessary to have in place measures to prevent their accidental release. These measures may be applied at national borders by prohibiting or controlling the importation of specified aquatic animal pathogens or pathological material, which may contain them.

Competent Authorities should not require sanitary measures for biological samples preserved for diagnostic applications that are treated in such a manner as to inactivate the pathogenic agent.

Article 5.10.2.

Importation of aquatic animal pathogens

The importation of a pathogen referred to in the Aquatic Code, whether in culture, in pathological material or in any other form, should be officially controlled by the Competent Authority to ensure appropriate safeguards are in place to manage the risk posed by the pathogen. The conditions should be appropriate to the risk posed by the pathogen and, in relation to air transport, the appropriate standards of the International Air Transport Association or other relevant transport associations concerning the packaging and transport of dangerous goods as outlined in Article 5.10.3. should apply.

When considering applications to import a pathogen referred to in the Aquatic Code, whether in culture, in pathological material or in any other form, from other countries, Competent Authorities should have regard to the nature of the material, the animal from which it is derived, the susceptibility of that animal to various diseases and the animal health situation of the country of origin. It may be advisable to require that material be pretreated before import to minimise the risk of inadvertent introduction of a pathogen referred to in the Aquatic Code.

Any material that does not satisfy the applied conditions should be rendered safe by the Aquatic Animal Health Service of the receiving country.

Article 5.10.3.

Packaging and documentation for transport

The safe transport of a pathogen referred to in the Aquatic Code, with respect to the pathogen, the handlers and the environment, is primarily dependent on proper packaging and it is the responsibility of the sender that this is done in accordance with current regulations.
1. **Basic triple packaging system**

The system consists of three layers as follows:

a) *Primary receptacle:* a labelled primary watertight, leak-proof receptacle containing the specimen. The receptacle is wrapped in enough absorbent material to absorb all fluid in case of breakage.

b) *Secondary receptacle:* a second durable, watertight, leak-proof receptacle to enclose and protect the primary receptacle(s). Several wrapped primary receptacles may be placed in one secondary receptacle. Sufficient additional absorbent material must be used to cushion multiple primary receptacles.

c) *Outer shipping package:* the secondary receptacle is placed in an outer shipping package, which protects it and its contents from outside influences such as physical damage, temperature fluctuations and water while in transit.

Ice or dry ice when used in a shipment must be placed outside the secondary receptacle. If wet ice is used, it should be in a leak-proof *container* and the outer package must also be leak-proof. The secondary receptacle must be secured within the outer package to prevent damage after the refrigerant has melted or dissipated.

Dry ice must *NOT* be placed inside the primary or secondary receptacle because of the risk of explosions. The outer package must permit the release of carbon dioxide gas if dry ice is used. IATA Packing Instruction 904 must be observed for packages containing dry ice.

2. **Documentation**

Specimen data forms, letters and other types of information that identify or describe the specimen and also identify the shipper and receiver should be taped to the outside of the secondary receptacle, together with a copy of the recipient's import permit.

**Article 5.10.4.**

Any sender of a pathogen referred to in the *Aquatic Code or pathological material* must ensure that the proposed receiver has obtained the necessary import licence referred to in Article 5.10.2.

**Article 5.10.5.**

1) Every consignment of a pathogen referred to in the *Aquatic Code or pathological material* should be notified in advance by the sender to the intended recipient, giving the following information:

a) exact nature of the sample and its packaging;

b) the number of packages sent and the marks and numbers enabling their identification;

c) date of despatch;

d) method of transport used for consignment of products (ship, aircraft, railway wagon or road vehicle).

2) The recipient should notify the sender of the receipt of each consignment of a pathogen referred to in the *Aquatic Code or pathological material* on its arrival.

3) When a consignment that has been notified by the sender fails to arrive by the anticipated date, the intended recipient should notify the *Competent Authority* of the receiving country and, at the same time, the sender in the country of origin, so that any necessary action can be taken for investigation to be made without delay.

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**NB:** FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2010.
CHAPTER 5.11.

MODEL HEALTH CERTIFICATES
FOR INTERNATIONAL TRADE
IN LIVE AQUATIC ANIMALS
AND PRODUCTS OF AQUATIC ANIMAL ORIGIN

Article 5.11.1.

Notes for guidance on the health certificates for international trade in live aquatic animals and products of aquatic animal origin

1. General
   Please complete the certificate on paper in capital letters. To confirm an option, mark the box with a cross (X). Ensure that no portion of certificate is left blank in a manner that would allow it to be amended. Non-applicable fields may be crossed out.

2. Part I. Details of dispatched consignment

| Box I.1. | Name and full address of the natural or legal entity dispatching the consignment. Information on telephone and fax numbers or e-mail address is recommended. |
| Box I.2. | The certificate reference number is the number used by the Competent Authority of the country to identify the certificate. |
| Box I.3. | Name of the Competent Authority. |
| Box I.4. | Name and full address of the natural or legal entity to whom the consignment is destined at the time the certificate is issued. |
| Box I.5. | Name of the country from which the live aquatic animals or gametes are being exported. For aquatic animal products, name the country(ies) where the finished products were produced, manufactured or packed. "ISO code" refers to the international standard two-letter code (ISO 3166-1 Alpha-2 Code) for a country produced by the International Organization for Standardization. |
| Box I.6. | Name of the zone or compartment of origin, if relevant, in part II of the certificate. |
| Box I.7. | Name of the country of destination. "ISO code" refers to the international standard two-letter code (ISO 3166-1 Alpha-2 Code) for a country produced by the International Organization for Standardization. |
| Box I.8. | Name of the zone or compartment of destination, if relevant, in part II of the certificate. |
| Box I.9. | Name and full address of the place(s) from which the live aquatic animals, gametes or aquatic animal products are being exported; and official approval or registration number when required. For live aquatic animals and gametes: the establishment(s) or place of capture. For products of aquatic animal origin: the premises from which the products are to be dispatched. |
| Box I.10. | Name of the place from which the live aquatic animals, gametes or aquatic animal products are being shipped (this will be a land, sea or airport). |
**Chapter 5.11.- Model health certificates for international trade in live aquatic animals and products of aquatic animal origin**

| Box I.11. | Date of departure. For live aquatic animals and gametes include the expected time of departure. |
| Box I.12. | Details of the means of transport. |
| Box I.13. | Identification of the means of transport at the time the certificate is issued: for air transport, the flight number; for maritime transport, the name of the vessel; for rail transport, the number of the train and the wagon and for road transport, the registration number of the road vehicle and the number of the trailer where used. |
| Box I.15. | CITES permit number(s) if the commodity concerns species listed in the Convention on International Trade in Endangered Species of Wild Fauna and Flora. |
| Box I.16. | Describe the commodity or use the titles as they appear in the Harmonised System of the World Customs Organization. |
| Box I.17. | Heading or HS Code of the Harmonized System set up by the World Customs Organization. |
| Box I.18. | Total quantity or weight of the commodity. |
| Box I.19. | Total number of containers in which they are being transported. For aquatic animal products give the total number of packages. |
| Box I.20. | Identify the containers/seal numbers where required. |
| Box I.21. | Intended use of the imported live aquatic animals or aquatic animal products. |
| Box I.22. | Ornamental: applies to live aquatic animals kept for companionship or enjoyment. |

**Breeding**: applies to gametes and broodstock.

**Grow out**: applies to live aquatic animals, aquatic eggs and aquatic larvae requiring time in culture.

**Slaughter**: applies to live aquatic animals for slaughter.

**Restocking**: applies to live aquatic animals for the purpose of rebuilding stocks.

**Competition/display**: applies to live aquatic animals used for competition or display purposes.

**Human consumption**: applies to live aquatic animals (without further aquaculture involved) or aquatic animals products intended for human consumption.

**Aquatic animal feed**: means any product of animal origin (single or multiple), whether processed, semi-processed or raw, that is intended to be fed to aquatic animals.

**Further processing**: applies to products of aquatic animal origin that have to be further processed before being suitable for end use.

**Other technical use**: applies to aquatic animal products not intended for human or aquatic animal consumption. These include aquatic animal products that are intended for use in the pharmaceutical, medical, cosmetic and other industries. Such products may be subjected to extensive further processing.

**Technical use in live aquatic animals**: applies to aquatic animal products used in live aquatic animals, e.g. to stimulate ovulation.
## Part II. Zoosanitary information

<table>
<thead>
<tr>
<th>Box II.</th>
<th>Complete this part in accordance with the requirements agreed between the Competent Authorities of the importing and exporting countries in accordance with the recommendations in the Aquatic Code.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Box II.a.</td>
<td>Reference number: see box I.2.</td>
</tr>
<tr>
<td>Certifying Official</td>
<td>Name, address, official position, date of signature and official stamp of the Competent Authority.</td>
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</table>
### Article 5.11.2.

**Model health certificate for international trade in live aquatic animals and gametes**

**COUNTRY:**

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<thead>
<tr>
<th>Part I. Details of dispatched consignment</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>I.1. Consignor: Name: Address:</td>
<td>I.2. Certificate reference number:</td>
</tr>
<tr>
<td>I.4. Consignee: Name: Address:</td>
<td>I.3. Competent Authority:</td>
</tr>
<tr>
<td>I.9. Place of origin: Name: Address:</td>
<td></td>
</tr>
<tr>
<td>I.10. Place of shipment:</td>
<td>I.11. Date of departure:</td>
</tr>
<tr>
<td>Aeroplane □</td>
<td>Ship □ Railway wagon □</td>
</tr>
<tr>
<td>Road vehicle □</td>
<td>Other □</td>
</tr>
<tr>
<td>Identification:</td>
<td></td>
</tr>
<tr>
<td>I.17. Total quantity:</td>
<td></td>
</tr>
<tr>
<td>I.18.</td>
<td>I.19. Total number of packages:</td>
</tr>
<tr>
<td>I.20. Identification of container/seal number:</td>
<td>I.21. Type of packaging:</td>
</tr>
<tr>
<td>I.22. Commodities intended for use as:</td>
<td></td>
</tr>
<tr>
<td>Breeding □</td>
<td>Grow out □</td>
</tr>
<tr>
<td>Slaughter □</td>
<td>Restocking □</td>
</tr>
<tr>
<td>Ornamental □</td>
<td>Competition/exhibition □</td>
</tr>
<tr>
<td>Other □ If other, specify.</td>
<td></td>
</tr>
<tr>
<td>I.23. For import or admission:</td>
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</tr>
<tr>
<td>Definitive import □ Re-entry □ Temporary admission □</td>
<td></td>
</tr>
<tr>
<td>I.24. Identification of commodities:</td>
<td></td>
</tr>
<tr>
<td>Amphibian □</td>
<td>Crustacean □</td>
</tr>
<tr>
<td>Fish □</td>
<td>Mollusc □</td>
</tr>
<tr>
<td>Wild stock □</td>
<td>Cultured stock □</td>
</tr>
<tr>
<td>Species (scientific name):</td>
<td>Age*:</td>
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<tr>
<td>Identification system*:</td>
<td>Batch number*:</td>
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<tr>
<td>Sex*:</td>
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* Optional. ** If referenced in Part II.
### COUNTRY:

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<th>Part II. Zoosanitary information</th>
<th>II.a. Certificate reference number:</th>
</tr>
</thead>
<tbody>
<tr>
<td>The undersigned Certifying Official certifies that the animal(s)/gametes described above satisfy(ies) the following requirements:</td>
<td></td>
</tr>
</tbody>
</table>

Certifying Official:

Name and address (in capital letters):

Official position:

Date:

Signature:

Stamp:
### Model health certificate for international trade in products of aquatic animal origin

#### COUNTRY:

<table>
<thead>
<tr>
<th>Part I: Details of dispatched consignment</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>I.1. Consignor: Name: Address:</td>
<td>1.2. Certificate reference number:</td>
</tr>
<tr>
<td>I.4. Consignee: Name: Address:</td>
<td>1.3. Competent Authority:</td>
</tr>
<tr>
<td>I.9. Place of origin: Name: Address:</td>
<td>1.10. Place of shipment:</td>
</tr>
<tr>
<td>I.12. Means of transport:</td>
<td>1.11. Date of departure:</td>
</tr>
<tr>
<td>Aeroplane □ Ship □ Railway wagon □</td>
<td>1.13. Expected border post:</td>
</tr>
<tr>
<td>Road vehicle □ Other □</td>
<td>1.14. CITES permit No(s)**:</td>
</tr>
<tr>
<td>Identification:</td>
<td>1.15. Description of commodity:</td>
</tr>
<tr>
<td>I.18. Temperature of the product:</td>
<td>1.19. Total number of packages:</td>
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<tr>
<td>Ambient □ Chilled □ Frozen □</td>
<td>1.20. Identification of container/seal number:</td>
</tr>
<tr>
<td>I.19. Total number of packages:</td>
<td>1.21. Type of packages:</td>
</tr>
<tr>
<td>I.22. Commodities intended for use as:</td>
<td></td>
</tr>
<tr>
<td>Human consumption □</td>
<td>Aquatic animal feed □</td>
</tr>
<tr>
<td>Further processing □</td>
<td>Other technical use □</td>
</tr>
<tr>
<td>Other □ If other, specify.</td>
<td>Technical use in aquatic animals □ If technical use, specify:</td>
</tr>
<tr>
<td>I.24. Identification of commodities:</td>
<td></td>
</tr>
<tr>
<td>Amphibian □</td>
<td>Crustacean □</td>
</tr>
<tr>
<td>Fish □</td>
<td>Mollusc □</td>
</tr>
<tr>
<td>Wild stock □</td>
<td>Cultured stock □</td>
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<td>Species (scientific name):</td>
<td>Approval number of establishments:</td>
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<tr>
<td>Lot ID/date code:</td>
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</tr>
</tbody>
</table>

* Optional.

** If referenced in Part II.
Chapter 5.11.- Model health certificates for international trade in live aquatic animals and products of aquatic animal origin

**COUNTRY:**

<table>
<thead>
<tr>
<th>Part II. Zoosanitary information</th>
<th>II.a. Certificate reference number:</th>
</tr>
</thead>
</table>

The undersigned Certifying Official certifies that the product(s) of aquatic animal origin described above satisfy(ies) the following requirements:

Certifying Official:

Name and address (in capital letters): Official position:

Date: Signature:

Stamp:

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2010.
SECTION 6.

ANTIMICROBIAL USE IN AQUATIC ANIMALS

CHAPTER 6.1.

INTRODUCTION TO THE RECOMMENDATIONS FOR CONTROLLING ANTIMICROBIAL RESISTANCE

Article 6.1.1.

Objectives

The purpose of this section is to provide guidance for Member Countries to appropriately address the selection and dissemination of resistant microorganisms and antimicrobial resistance determinants from the use of antimicrobial agents in aquatic animals.

Antimicrobial agents are essential for human and animal health and welfare. The OIE recognises the need for access to antimicrobial agents in veterinary medicine: antimicrobial agents are essential for treating and controlling infectious diseases in aquatic animals. The OIE therefore considers that ensuring continued access to effective antimicrobial agents is important.

The OIE recognises that antimicrobial resistance is a global public and animal health concern that is influenced by the usage of antimicrobial agents in humans, animals and elsewhere. Those working in the human, animal and plant sectors have a shared responsibility to address the risk factors for the selection and dissemination of antimicrobial resistance. Arising from its mandate for the protection of animal health and food safety, the OIE developed these chapters to provide guidance to Member Countries in regard to risks in the animal sector.

The application of risk assessment and risk management measures should be based on relevant international standards on risk analysis and supported by sound data and information when available. The guidance provided in these chapters should be consulted as part of the standard approach to reduce the risk associated with the selection and dissemination of antimicrobial resistant microorganisms and antimicrobial resistance determinants.

NB: FIRST ADOPTED IN 2010; MOST RECENT UPDATE ADOPTED IN 2011.
CHAPTER 6.2.

PRINCIPLES FOR RESPONSIBLE AND PRUDENT USE OF ANTIMICROBIAL AGENTS IN AQUATIC ANIMALS

Article 6.2.1.

Purpose

These principles provide guidance for the responsible and prudent use of antimicrobial agents in aquatic animals, with the aim of protecting both animal and human health. The Competent Authorities responsible for the registration and marketing authorisation of products and the control of all organisations involved in the production, distribution and use of antimicrobial agents have specific obligations.

Article 6.2.2.

Objectives of responsible and prudent use

Responsible and prudent use includes a set of practical measures and recommendations intended to reduce the risk associated with the selection and dissemination of antimicrobial resistant microorganisms and antimicrobial resistance determinants in aquatic animal production to:

1) maintain the efficacy of antimicrobial agents both for veterinary and human medicine and to ensure the rational use of antimicrobials in aquatic animals with the purpose of optimising both their efficacy and safety;
2) comply with the ethical obligation and economic need to keep aquatic animals in good health;
3) prevent or reduce the transfer of both resistant microorganisms and resistance determinants from aquatic animals to humans and terrestrial animals;
4) prevent antimicrobial residues that exceed the established maximum residue limit (MRL) occurring in the food.

Article 6.2.3.

Definition

Pharmacovigilance of antimicrobial agent: means the detection and investigation of the effects of the use of these products, mainly aimed at safety and efficacy in aquatic animals and safety in people exposed to the products.

Article 6.2.4.

Responsibilities of Competent Authorities

The Competent Authorities responsible for granting marketing authorisation for antimicrobial agents have a significant role in specifying the terms of the authorisation and in providing the appropriate information to the veterinarian or other aquatic animal health professional through product labelling and/or by other means, in support of prudent use of antimicrobial agents in aquatic animals.

It is the responsibility of Competent Authorities to develop up-to-date guidelines on data requirements for evaluation of antimicrobial agent applications.

Competent Authorities in cooperation with animal and public health professionals should adopt a proactive approach to promote prudent use of antimicrobial agents in aquatic animals as an element of a comprehensive strategy for the containment of antimicrobial resistance.

Elements of a comprehensive strategy should include good animal husbandry practices, vaccination policies and development of animal health care at the farm level, and consultation with a veterinarian or other aquatic animal health professional.
Chapter 6.2. - Principles for responsible and prudent use of antimicrobial agents in aquatic animals

professional, all of which should contribute to reduction of the prevalence of animal disease requiring antimicrobial treatment.

Competent Authorities should expeditiously grant marketing authorisations when criteria of quality, efficacy and safety are met.

The examination of marketing authorisation applications should include an assessment of the risks to animals, humans and the environment resulting from the use of antimicrobial agents in aquatic animals. The evaluation should focus on each individual antimicrobial agent and take into consideration the class of antimicrobials to which the particular active substance belongs. The safety evaluation should include consideration of the potential impact of the proposed use in aquatic animals on human health, including the human health impact of antimicrobial resistance developing in microorganisms found in aquatic animals. An assessment of the impact of the proposed use on the environment should be conducted.

Competent Authorities should aim to ensure that advertising of antimicrobial agents complies with relevant legislation and marketing authorisations granted and discourage direct advertising other than to those legally entitled to prescribe the antimicrobial agent.

Information collected through pharmacovigilance programmes, including on lack of efficacy, should form part of the Competent Authority’s comprehensive strategy to minimise antimicrobial resistance.

Competent Authorities should disseminate, to veterinarians or other aquatic animal health professionals, information on trends in antimicrobial resistance collected during surveillance programmes and should monitor the performance of susceptibility testing laboratories.

Competent Authorities and stakeholders should work together to provide for effective procedures for the safe collection and destruction of unused or out-of-date antimicrobial agents.

Article 6.2.5.

Responsibilities of the veterinary pharmaceutical industry

The veterinary pharmaceutical industry has responsibilities for providing information requested by Competent Authorities on the quality, efficacy and safety of antimicrobial agents. The responsibilities of the veterinary pharmaceutical industry cover pre- and post-marketing phases, including manufacturing, sale, importation, labelling, advertising and pharmacovigilance.

The veterinary pharmaceutical industry has the responsibility to provide the Competent Authority with the information necessary to evaluate the amount of antimicrobial agents marketed. The veterinary pharmaceutical industry should ensure that the advertising of antimicrobial agents directly to the aquatic animal producer is discouraged.

Article 6.2.6.

Responsibilities of wholesale and retail distributors

Distributors should ensure that their activities are in compliance with the relevant legislation.

Distributors should ensure that information for the appropriate use and disposal of the antimicrobial agent accompany all distributed products and should also be responsible for maintaining and disposing of the product in accordance with the manufacturer recommendations.

Article 6.2.7.

Responsibilities of veterinarians and other aquatic animal health professionals

Responsibilities of veterinarians or other aquatic animal health professionals include identifying, preventing and treating aquatic animal diseases, as well as the promotion of sound animal husbandry methods, hygiene procedures, vaccination and other alternative strategies to minimise the need for antimicrobial use in aquatic animals.
Veterinarians or other aquatic animal health professionals authorised to prescribe veterinary medicines should only prescribe, dispense or administer a specific course of treatment with an antimicrobial agent for aquatic animals under their care.

The responsibilities of veterinarians or other aquatic animal health professionals are to carry out a thorough clinical assessment of the aquatic animal(s), including as appropriate: clinical examination, post-mortem examination, bacteriology with culture and sensitivity, and other laboratory tests to arrive at the most definitive diagnosis possible before initiating a specific course of treatment with an antimicrobial agent. Evaluation of environmental factors and husbandry at the production site (e.g. water quality) should be considered as potential primary factors leading to infection and should be addressed prior to prescribing a course of antimicrobial agent treatment.

If therapy with an antimicrobial agent is deemed necessary it should be initiated as soon as possible. The selection of the agent should be based on the knowledge and experience of the veterinarian or other aquatic animal health professional authorised to prescribe veterinary medicines.

As soon as possible, susceptibility testing of the target microorganism should be used to confirm the choice of treatment. Results of all susceptibility tests should be retained and should be available to the Competent Authority.

The veterinarian or other aquatic animal health professional authorised to prescribe veterinary medicines should indicate precisely to the aquatic animal producer the treatment regime, including the dose, the treatment intervals, the duration of the treatment, the withdrawal period and the amount of antimicrobial agents to be delivered, depending on the dosage and the number of aquatic animals to be treated.

The use of antimicrobial agents extra-label/off-label may be permitted in appropriate circumstances in conformity with the relevant legislation.

Records on the use of antimicrobial agents should be kept in conformity with the relevant legislation. Veterinarians or aquatic animal health professionals should also periodically review farm records on the use of the antimicrobial agents to ensure compliance with their directions and use these records to evaluate the efficacy of treatment regimens. Suspected adverse reactions, including a lack of efficacy, should be reported to the Competent Authority. Associated susceptibility data should accompany the report of lack of efficacy.

Article 6.2.8.

Responsibilities of aquatic animal producers

Aquatic animal producers should implement health programmes on their farms in order to promote aquatic animal health and food safety. This can be done through adequate planning of culture strategies to maintain aquatic animal health through biosecurity programmes, husbandry, nutrition, vaccination, maintenance of good water quality, etc.

Aquatic animal producers should use antimicrobial agents only on the prescription of a veterinarian or other aquatic animal health professional authorised to prescribe veterinary medicines, and follow directions on the dosage, method of application, and withdrawal period.

Aquatic animal producers should ensure that antimicrobial agents are properly stored, handled, and disposed.

Aquatic animal producers should keep adequate records of antimicrobial agents used, bacteriological and susceptibility tests, and make such records available to the veterinarian or other aquatic animal health professional.

Aquatic animal producers should inform the veterinarian or other aquatic animal health professional of recurrent disease problems and lack of efficacy of antimicrobial agent treatment regimes.

Article 6.2.9.

Training of users of antimicrobial agents

The training of users of antimicrobial agents should involve all the relevant organisations, such as relevant regulatory authorities, pharmaceutical industry, veterinary schools, research institutes, and veterinary professional organisations and other approved users such as aquatic animal owners.
Research

To address the significant lack of information for numerous species of aquatic animals, the relevant regulatory authorities and other stakeholders should encourage public-funded and industry-funded research.

NB: FIRST ADOPTED IN 2011.
CHAPTER 6.3.

MONITORING OF THE QUANTITIES AND USAGE PATTERNS OF ANTIMICROBIAL AGENTS USED IN AQUATIC ANIMALS

Article 6.3.1.

Purpose

The purpose of these recommendations is to describe approaches to the monitoring of quantities of antimicrobial agents used in aquatic animals, including species reared for food and ornamental purposes.

These recommendations are intended for use in the collection of objective and quantitative information to evaluate usage patterns by antimicrobial class, route of administration and aquatic animal species in order to evaluate exposure of microorganisms to antimicrobial agents.

The collection of data on the use of antimicrobial agents in aquaculture may be constrained in some countries by the lack of available resources, lack of accurately labelled products, poorly documented distribution channels and lack of professional consultation or supervision. This chapter may therefore be seen as indicating the direction in which countries should develop with regard to collecting data and information on the use of antimicrobial agents in aquatic animals.

Article 6.3.2.

Objectives

The information provided in these recommendations is essential for conducting risk analyses and for planning purposes. This information can be helpful in interpreting antimicrobial resistance surveillance data and can assist in the ability to respond to problems of antimicrobial resistance in a precise and targeted way. The continued collection of this basic information would help identify trends in the use of antimicrobial agents in aquatic animals and the potential association with antimicrobial resistance in aquatic animal bacteria, including potentially zoonotic bacteria. This information may also assist in risk management when evaluating the effectiveness of efforts to ensure responsible and prudent use and mitigation strategies and indicate where alteration of prescribing practices for antimicrobial agents in aquatic animals might be appropriate. The publication of these data and their interpretation is important to ensure transparency and to allow all interested parties to assess trends, to perform risk assessments and for risk communication purposes.

Article 6.3.3.

Development and standardisation of monitoring systems for antimicrobial agents

Competent Authorities may, for reasons of cost and administrative efficiency, collect medical, agricultural, aquacultural and other antimicrobial agents use data in a single programme. Where livestock and aquatic animal industries are under multiple authorities in a single country, collaboration between the authorities to develop a coordinated monitoring system is necessary to facilitate the collection of data. Additionally, a consolidated programme would facilitate the comparison of aquatic animal use data with human use data necessary for a comprehensive risk analysis.

Systems to monitor usage of antimicrobial agents may consist of the following elements:
Chapter 6.3.- Monitoring of the quantities and usage patterns of antimicrobial agents used in aquatic animals

1. Sources of data on antimicrobial agents
   a) Basic sources
      Data from basic sources may include general information without specific attribution (such as, weight, quantity and class of antimicrobial agents).
      Sources of data will vary from country to country. Such sources may include customs, import, export, manufacturing and sales data.
   b) Direct sources
      Data from direct sources may include more specific information (such as target aquatic animal species, route of administration and active ingredient).
      Data from veterinary medicinal product registration authorities, manufacturers, wholesalers, retailers, feed stores and feed mills might be useful sources. A possible mechanism for the collection of this information is to make the provision of appropriate information by veterinary antimicrobial manufacturers to the registration authority one of the requirements of marketing authorisation (registration of the antimicrobial agent).
   c) End-use sources
      Data from end-use sources has the advantage of providing more detailed information on the type and purpose of use and can be complimentary to the other sources.
      End-use sources of data may include veterinarians, aquatic animal health professionals and aquatic animal producers. End-use sources may be useful when more accurate and locally specific information is needed (such as extra-/off-label use).
      Collection of this type of information can be resource intensive; therefore, periodic collection of this type of information may be sufficient. Data collection should be targeted to the most relevant period of use.
      In some countries end-use sources may be the only practical source of information.
   d) Other sources
      Pharmaceutical industry associations and aquatic animal producer associations, veterinary and allied health professional associations, and other stakeholders with indirect knowledge of the quantities of antimicrobial agents used may be another source of this information.
      Non-conventional sources including Internet sales data related to antimicrobial agents may be collected where available. Internet sales data may be particularly useful with respect to ornamental species.

2. Elements for data collection and reporting
   a) Basic data to be collected should include:
      i) the absolute amount in kilograms of the active ingredient of the antimicrobial agent(s) used per year, divided into antimicrobial class/subclass;
      for active ingredients present in the form of compounds or derivatives, the mass of active entity of the molecule should be recorded; for antimicrobial agents expressed in International Units, the calculation required to convert these units to mass of active entity should be stated; it may be possible to estimate total usage by collecting sales data, prescribing data, manufacturing data, export/import data or any combination of these;
      ii) the total number of aquatic animals treated and their weight in kilograms.
   b) Additional data may be collected to further categorise the exposure of microorganisms to antimicrobial agents and may include:
      i) species of fish, crustaceans, molluscs or amphibians treated;
      ii) purpose e.g. aquatic animals for human consumption, use as ornamental species and baitfish;
      iii) route of administration (medicated feed, bath treatment, parenteral delivery) and the method used to calculate the dose (biomass of aquatic animals, volume of water treated);
      iv) indication for use.
      The antimicrobial agents/classes/sub-classes to be included in data reporting should be based on current known mechanisms of antimicrobial activity / antimicrobial resistance mechanism.
      Nomenclature of antimicrobial agents should comply with international standards where available.
      When making information publicly available, the Competent Authority should ensure confidentiality and anonymity of individual enterprises.
Chapter 6.3.- Monitoring of the quantities and usage patterns of antimicrobial agents used in aquatic animals

3. Considerations for data collection

Antimicrobial usage data may be collected on a routine basis and/or at a specific point in time depending on availability of resources and/or the need to monitor usage of antimicrobial agents or address a specific antimicrobial resistance problem.

Registration of products with labelling that accurately reflects the intended use of the antimicrobial agent will facilitate collection of information on the quantities and usage patterns.

Collection, storage and processing of data from end-use sources requires careful design but should have the advantage of producing accurate and targeted information.

Article 6.3.4.

Elements for interpretation of data on the use of antimicrobial agents

When available, the following information may support the interpretation of antimicrobial usage data and further characterisation of exposure pathways:

1) type of aquaculture system (extensive or intensive, ponds or tanks, flow-through or recirculating, hatchery or grow-out, integrated system);
2) animal movements (transfer between facilities or from wild to the facility, grading);
3) species, life stage, and/or stage of the production cycle;
4) environmental and culture parameters (seasonality, temperature, salinity, pH);
5) geographical location, specific rearing units;
6) weight/biomass, dosage regimes and duration of treatment with antimicrobial agents;
7) basis for treatment (historical, empirical, clinical, clinical with laboratory confirmation and sensitivity testing).

Factors such as the number/percentage of animals/culture units treated, treatment regimens, type of use and route of administration are key elements to consider for risk assessment.

When comparing use of antimicrobial agents over time, changes in size and composition of animal populations should also be taken into account.

Regarding data coming from end-user sources, analysis of the use of antimicrobial agents may be possible at the regional, local or farm level, and at the level of the individual veterinarian or other aquatic animal health professional.

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NB: FIRST ADOPTED IN 2012.
CHAPTER 6.4.

DEVELOPMENT AND HARMONISATION OF NATIONAL ANTIMICROBIAL RESISTANCE SURVEILLANCE AND MONITORING PROGRAMMES FOR AQUATIC ANIMALS

Article 6.4.1.

Purpose

This chapter provides criteria relevant to aquatic animals and aquatic animal products intended for human consumption for:

1) the development of national antimicrobial resistance surveillance and monitoring programmes and
2) the harmonisation of existing national antimicrobial resistance surveillance and monitoring programmes.

Article 6.4.2.

Objective of surveillance and monitoring programmes

Competent Authorities should conduct active antimicrobial resistance surveillance and monitoring programmes for aquatic animals.

Surveillance and monitoring of antimicrobial resistance is necessary to:

1) establish baseline data on the prevalence of antimicrobial resistant microorganisms and determinants;
2) collect information on antimicrobial resistance trends in relevant microorganisms;
3) explore the potential relationship between antimicrobial resistance in aquatic animal microorganisms and the use of antimicrobial agents;
4) detect the emergence of antimicrobial resistance mechanisms;
5) conduct risk analyses as relevant to aquatic animal and human health;
6) provide recommendations on human health and aquatic animal health policies and programmes;
7) provide information to facilitate prudent use, including guidance for professionals prescribing the use of antimicrobial agents in aquatic animals.

Cooperation at a regional level between countries conducting antimicrobial resistance surveillance should be encouraged.

The findings of surveillance and monitoring programmes should be shared at the regional and international level to maximise understanding of the global risks to aquatic animal health and human health. The publication of these data and their interpretation is important to ensure transparency and to allow all interested parties to assess trends, to perform risk assessments and for risk communication purposes.

Article 6.4.3.

General considerations for the design of surveillance and monitoring programmes

Surveillance of antimicrobial resistance at targeted intervals or ongoing monitoring of the prevalence of resistance in microorganisms from aquatic animals, aquatic animal products intended for human consumption, and humans constitutes a critical part of aquatic animal health and public health strategies aimed at limiting the spread of antimicrobial resistance and optimising the choice of antimicrobial agents used in therapy.

For aquaculture it is important to conduct surveillance and monitoring of microorganisms that infect aquatic animals and microorganisms, including human pathogens, present on food derived from aquatic animals.
Design of surveillance and monitoring programmes for antimicrobial susceptibility of microorganisms that infect aquatic animals

An important consideration for the design of surveillance and monitoring programmes for antimicrobial susceptibility of microorganisms that infect aquatic animals is the lack of standardised and validated antimicrobial testing methods for a significant number of bacterial species of aquatic importance. When validated methods are available they should be used. Any deviations from standard methodology should always be clearly reported. For tests performed on bacterial species for which standard methods have not been developed full details of the methods used should be provided.

A preliminary requirement for the development of a surveillance and monitoring programme may be the identification and prioritisation of bacteria isolated from aquatic animals for methods development.

1. Selection of microorganisms
   Information on the occurrence of antimicrobial resistance in microorganisms that infect aquatic animals should be derived from regular monitoring of isolates obtained from diagnostic laboratories. These isolates should have been identified as primary causal agents of significant disease epizootics in aquatic animals.
   It is important that monitoring programmes focus on microorganisms that are associated with the commonly encountered infections of the major aquatic species farmed in the region / local growing area.
   Selection should be designed to minimise bias resulting from over representation of isolates obtained from severe epizootics or epizootics associated with therapeutic failures.
   Microorganisms belonging to a specific species or group may be selected for intensive study in order to provide information on a particular problem.

2. Methods used to analyse microorganism susceptibility to antimicrobial agents
   Participating laboratories may perform disc diffusion, minimum inhibitory concentration (MIC) or other susceptibility tests to monitor frequencies of resistance. Protocols that have been standardised internationally and validated for application to the study of microorganisms isolated from aquatic animals should always be used.

3. Requirements for laboratories involved in monitoring resistance
   Laboratories involved in national or regional monitoring of antimicrobial resistance should be of sufficient capability and have relevant expertise to comply with all the quality control requirements of the standardised test protocols. They should also be capable of participating in all necessary inter-laboratory calibration studies and method standardisation trials.

4. Choice of antimicrobial agents
   Representatives of all major classes of antimicrobial agents used to treat disease in aquatic animal species should be included in susceptibility testing.

5. Reporting of results
   The results of surveillance and monitoring programmes, including susceptibility data, should be published and made available for use by relevant stakeholders. Both primary quantitative data and the interpretive criteria used should be reported.

6. Surveillance and monitoring for epidemiological purposes
   For epidemiological surveillance purposes, use of the epidemiological cut-off value (also referred to as microbiological breakpoint), which is based on the distribution of MICs or inhibition zone diameters of the specific microbial species tested, is preferred.
   When reporting interpretations made by application of epidemiological cut-off values, the resultant categories should be referred to as wild type (WT) or non-wild type (NWT). When interpretations are made by the application of breakpoints the resultant categories should be referred to as sensitive, intermediate or resistant.
   For microbial species and antimicrobial agent combinations, where internationally agreed epidemiological cut-off values have not been set, laboratories may establish their own laboratory-specific values provided the methods they use are clearly reported.

7. Surveillance and monitoring for clinical purposes
   The application of clinical breakpoints may be appropriate when the aim of the programme is to provide information to facilitate prudent use, including guidance for professionals in prescribing antimicrobial agents in aquatic animals.
Selecting antimicrobial agents for therapeutic administration on the basis of information gained from the application of validated clinical breakpoints to antimicrobial susceptibility test data for microorganisms isolated from aquatic animals is an important element in the prudent use of these agents.

Use of these clinical breakpoints allows microorganisms to be identified as unlikely to respond to the in vivo concentrations of antimicrobial agents achieved by a given standard therapeutic regime. In order to facilitate the development of these breakpoints, data is required that allows clinical correlation to be completed. For this purpose, where possible, data that relates in vitro susceptibility of isolates to the clinical outcome of treatments with specified dose regimes under specific environmental conditions should be collected and reported.

Valuable information with respect to setting clinical breakpoints can be gained from situations where therapeutic failure is reported. The Competent Authority should include, in a surveillance and monitoring programme, systems for capturing details of failed treatments and the laboratory susceptibility test of the microorganisms involved.

Article 6.4.5.

Design of surveillance and monitoring programmes for microorganisms in or on aquatic animal products intended for human consumption

For details of the sampling protocols and analytical procedures required for surveillance and monitoring programmes for antimicrobial resistance in microorganisms present in aquatic animal products intended for human consumption, Chapter 6.7. of the OIE Terrestrial Animal Health Code should be consulted.

It is important to note that the word ‘commensal’ as used in Chapter 6.7. of the OIE Terrestrial Animal Health Code has less relevance due to the transient nature of the intestinal microflora of aquatic animals. The inclusion of intestinal microflora in surveillance and monitoring programmes should only be considered when there is evidence that these are resident for sufficient time to be a risk factor affected by antimicrobial agents.

When designing a sampling programme it is important to consider that contamination of aquatic animal products with resistant microorganisms that are capable of infecting humans may arise from sources other than the aquatic animal. All sources of contamination should be taken into account, for example entry of raw manure into the aquatic environment. The number of such microorganisms associated with aquatic animals is much less than that found in terrestrial animals. However the following species should be included, as a minimum, in a surveillance and monitoring programme:

1) Salmonella spp.;
2) Vibrio parahaemolyticus;
3) Listeria monocytogenes.

NB: FIRST ADOPTED IN 2012.
CHAPTER 6.5.

RISK ANALYSIS FOR ANTIMICROBIAL RESISTANCE ARISING FROM THE USE OF ANTIMICROBIAL AGENTS IN AQUATIC ANIMALS

Article 6.5.1.

Recommendations for analysing the risks to aquatic animal health and human health from antimicrobial resistant microorganisms of aquatic animal origin

1. Introduction
Antimicrobial resistance is a naturally occurring phenomenon influenced by many factors. However, problems related to antimicrobial resistance are inherently related to antimicrobial agent use in any environment, including human and non-human uses.

Antimicrobial resistance associated with the use of antimicrobial agents for therapeutic and non-therapeutic purposes has led to the selection and dissemination of antimicrobial resistant microorganisms, with a resulting loss of therapeutic efficacy in animal and human medicine of antimicrobial agents.

2. Objective
For the purposes of this chapter, the principal aim of risk analysis is to provide Member Countries with a transparent, objective and scientifically defensible method of assessing and managing the human and aquatic animal health risks associated with the selection and dissemination of resistance arising from the use of antimicrobial agents in aquatic animals.

Guidance on the issue of foodborne antimicrobial resistance related to the non-human use of antimicrobial agents is covered by the Codex Guidelines for Risk Analysis of Foodborne Antimicrobial Resistance (CAC/GL77-2011).

3. Definitions
For the purposes of this chapter, the hazard is the resistant microorganism or resistance determinant that emerges as a result of the use of a specific antimicrobial agent in aquatic animals. This definition reflects the potential for resistant microorganisms to cause adverse health effects, as well as the potential for horizontal transfer of genetic determinants between microorganisms. The conditions under which the hazard might produce adverse consequences include any scenarios through which humans or aquatic animals could become exposed to an antimicrobial resistant pathogen, fall ill and then be treated with an antimicrobial agent that is no longer effective.

For the purposes of this chapter, risk to aquatic animal health relates to the infection of aquatic animals with microorganisms in which resistance has emerged due to antimicrobial agent usage in aquaculture, and resulting in the loss of benefit of antimicrobial therapy used to manage aquatic animal diseases.

For the purposes of this chapter, risk to human health relates to the infection of humans with microorganisms in which resistance has emerged due to antimicrobial agent usage in aquatic animals, and resulting in the loss of benefit of antimicrobial therapy used to manage the human infection.

4. The risk analysis process
The components of risk analysis described in this chapter are hazard identification, risk assessment, risk management and risk communication.

The chapter includes factors to be considered at various steps of the risk analysis process. These factors are not intended to be exhaustive and not all elements may be applicable in all situations.

5. Risk assessment
The assessment of the risk to human and aquatic animal health from antimicrobial resistant microorganisms resulting from the use of antimicrobial agents in aquatic animals should examine:

a) the likelihood of emergence of resistant microorganisms arising from the use of an antimicrobial agent, or more particularly dissemination of the resistance determinants if transmission is possible between microorganisms;
b) all pathways and their contribution to the likelihood of humans and aquatic animals being exposed to these resistant microorganisms or resistance determinants;

c) the consequences of exposure in terms of risks to human and aquatic animal health.

The general principles of risk assessment as defined in Article 2.1.3. apply equally to both qualitative and quantitative risk assessment.

Article 6.5.2.

Special considerations for conducting antimicrobial resistance risk analysis in aquaculture

1. Introduction

Antimicrobial resistance (AMR) risk analysis in aquaculture is challenged by a variety of factors that impact both risk assessment and risk management, including the diversity of aquaculture, relative lack of methods for culture and antimicrobial susceptibility testing (AST), relative lack of information on use of drugs, and potential for the development of a reservoir of resistant microorganisms and resistance determinants with a potential for horizontal transmission.

Nevertheless, the fundamental principles of risk analysis (risk assessment, risk management, risk communication) provide a framework just as valuable for aquaculture as for terrestrial animal production.

2. Data needs

Special care is required in the design of data collection programmes for risk assessment to take account of possible confounding factors.

Because many types of aquaculture operations (in particular, open systems) intersect with terrestrial animal production and human environments, it is especially important to clearly identify the risk to be assessed. The selection and dissemination of resistant microorganisms or resistant determinants may be associated with the use of antimicrobial agents on aquatic animals or it may be the result of antimicrobial use in nearby terrestrial animal production operations or the presence of antimicrobial agents in human waste water.

3. Diversity of aquaculture

The range of species under culture, the number and type of different culture systems, and the range of antimicrobial agents and their routes of administration impact elements of the risk assessment, particularly the entry assessment. Therefore, careful attention should be used when grouping seemingly similar sectors of the aquaculture industry.

Identification, selection and monitoring of risk management options are also influenced by the diversity of aquaculture.

4. Lack of standardised methods for antimicrobial susceptibility testing

Currently, standardised methods for antimicrobial susceptibility testing (AST) for many relevant aquaculture species are lacking resulting in inability to quantify specific risks. Standardised AST methods should be used where available; or when standardised methods are not available, well-described and scientifically sound approaches should be applied.

5. Lack of approved drugs

The small number of approved antimicrobial agents for use in aquaculture challenges risk analysis, both in terms of risk assessment and risk management.

The collection and use of thorough information on the types and quantities of antimicrobial agents that are in use in aquaculture and relevant to the risk assessment is important. In some circumstances legal extra or off-label and illegal uses may also need to be considered. See Chapter 6.3.

For risk management, the small number of approved drugs in combination with a range of regulatory and aquatic animal health infrastructure in countries engaged in aquaculture presents additional challenges. Risk management options should be practical and take into account the ability for enforcement and compliance.

For monitoring and surveillance programmes, a lack of approved drugs means systems for collection of data and information on the quantities of antimicrobial agents used may need to consider not only licensed distribution of approved drugs, but information on the use of unapproved drugs.
6. Potential for development of a reservoir (horizontal transmission)

Microorganisms inhabiting the environment represent the fundamental reservoir of resistant determinants in the biosphere. This reservoir represents the basic origin of all antimicrobial agent resistance determinants encountered in human and veterinary medicine. The frequency of resistance determinants in environmental microorganisms is maintained by intrinsic, non-anthropogenic factors; all human uses of antimicrobial agents, including in aquaculture, have the potential to increase the size of the reservoir.

There is a risk that the use of antimicrobial agents in aquaculture will result in a rise in the frequency of resistance determinants in the environmental microbiome. This may result in an increase in the frequency with which determinants are transferred to microorganisms capable of infecting humans, animals or aquatic animals. The assessment and management of this risk are extremely complex. The biological pathways both for the entry assessment and the exposure assessment are myriad and at present no specific guidelines can be offered.

Article 6.5.3.

Analysis of risks to human health

1. Definition of the risk

The infection of humans with microorganisms in which resistance has emerged due to antimicrobial agent usage in aquatic animals, and resulting in the loss of benefit of antimicrobial therapy used to manage the human infection.

2. Hazard

- Microorganisms that have acquired resistance (including multiple resistance) arising from the use of an antimicrobial agent in aquatic animals.
- Microorganisms having obtained a resistance determinant from other microorganisms which have acquired resistance arising from the use of an antimicrobial agent in aquatic animals.

The identification of the hazard should include consideration of the class or subclass of the antimicrobial agent. This definition should be read in conjunction with point 3 of Article 6.5.1.

3. Entry assessment

An entry assessment describes the biological pathways by which the use of a specific antimicrobial agent in aquatic animals leads to the entry of resistant microorganisms or resistance determinants into a particular environment. This assessment includes estimating qualitatively or quantitatively the probability of that complete process occurring. The entry assessment describes the probability of the entry of each of the hazards under each specified set of conditions with respect to amounts and timing.

The following factors should be considered in the entry assessment:
- species of aquatic animals treated with the antimicrobial agent(s) in question;
- aquaculture production system (intensive or extensive, net pens, tanks, raceways, ponds, other);
- number of aquatic animals treated, their age and their geographical distribution;
- prevalence of disease for which the antimicrobial agent is indicated or is used in the target aquatic animal population;
- data on trends in antimicrobial agent use and changes in aquaculture production systems;
- data on potential extra-label or off-label use;
- methods and routes of administration of the antimicrobial agent;
- dosage regimen (dose, dosing interval and duration of the treatment);
- pharmacokinetics and relevant pharmacodynamics of the antimicrobial agent;
- site and type of infection;
- development of resistant microorganisms;
- prevalence of pathogenic agents that are likely to develop resistance in an aquatic animal species;
- mechanisms and pathways of direct or indirect transfer of resistance;
- potential linkage of virulence attributes and resistance;
- cross-resistance or co-resistance with other antimicrobial agents;
data on trends and occurrence of resistant microorganisms obtained through surveillance of aquatic animals and aquatic animal products and waste products.

The following confounding factors should be considered in the entry assessment:

- resistant microorganisms or resistant determinants associated with aquatic animals or aquatic animal products that are a result of terrestrial contamination of the aquatic environment, feed contamination or contamination during post-harvest processing.

4. Exposure assessment

An exposure assessment describes the biological pathways necessary for exposure of humans to the resistant microorganisms or resistance determinants released from a given antimicrobial agent's use in aquatic animals, and estimates the probability of exposures occurring. The probability of exposure to the identified hazards is estimated for specified exposure conditions with respect to amounts, timing, frequency, duration of exposure, routes of exposure, and other characteristics of the human populations exposed.

The following factors should be considered in the exposure assessment:

- human demographics, including population subgroups, food consumption patterns, and traditions and cultural practices with respect to the preparation and storage of food;
- prevalence of resistant microorganisms in food at the point of consumption;
- microbial load in contaminated food at the point of consumption;
- environmental contamination with resistant microorganisms;
- transfer of resistant microorganisms and their resistance determinants between humans, aquatic animals, and the environment;
- measures taken for microbial decontamination of food;
- survival capacity and dissemination of resistant microorganisms during the food production process (including slaughtering, processing, storage, transportation and retailing);
- disposal practices for waste products and the likelihood for human exposure to resistant microorganisms or resistance determinants through those waste products;
- capacity of resistant microorganisms to become established in humans;
- human-to-human transmission of the microorganisms under consideration;
- capacity of resistant microorganisms to transfer resistance to human commensal microorganisms and zoonotic agents;
- amount and type of antimicrobial agents used to treat humans;
- pharmacokinetics, such as metabolism, bioavailability, distribution to the gastrointestinal flora;
- level of direct contact of workers in the aquaculture or processing industries to the antimicrobial resistant organisms.

5. Consequence assessment

A consequence assessment describes the relationship between specified exposures to resistant microorganisms or resistance determinants and the consequences of those exposures. A causal process should exist by which exposures produce adverse health or environmental consequences, which may in turn lead to socio-economic consequences. The consequence assessment describes the potential consequences of a given exposure and estimates the probability of them occurring.

The following factors should be considered in the consequence assessment:

- microbial dose and subsequent host response interactions;
- variation in susceptibility of exposed populations or subgroups of the population;
- variation and frequency of human health effects resulting from loss of efficacy of antimicrobial agents and associated costs (e.g. illness and hospitalisation);
- potential linkage of virulence attributes and resistance;
- changes in food consumption patterns due to loss of confidence in the safety of food products and any associated secondary risks;
- interference with antimicrobial therapy in humans;
- importance of the antimicrobial agent in animal health and human health (see OIE List of Antimicrobial Agents of Veterinary Importance and WHO List of Critically Important Antimicrobials);
- prevalence of resistance in human bacterial pathogens under consideration.
6. **Risk estimation**

A risk estimation integrates the results from the entry assessment, exposure assessment and consequence assessment to produce overall estimates of risks associated with the hazards. Thus, risk estimation takes into account the whole of the risk pathway from hazard identification to the unwanted consequences.

7. **Risk management**

*Risk management* consists of the steps described below.

a) **Risk evaluation**

Risk evaluation - the process of comparing the risk estimated in the *risk assessment* with the reduction in risk expected from the proposed *risk management* measures.

b) **Option evaluation**

A range of *risk management* options is available to minimise the emergence and dissemination of antimicrobial resistance and these include both regulatory and non-regulatory options, such as the development of codes of practice for the use of *antimicrobial agents* in *aquaculture*. *Risk management* decisions need to consider fully the implications of these different options for human health and *aquatic animal* health and welfare and also take into account economic considerations and any associated environmental issues. Effective control of *aquatic animal diseases* can have the dual benefits of reducing the risks to human health associated with both the bacterial pathogen under consideration and antimicrobial resistance.

c) **Implementation**

Risk managers should develop an implementation plan that describes how the decision will be implemented, by whom and when. *Competent Authorities* should ensure an appropriate regulatory framework and infrastructure.

d) **Monitoring and review**

*Risk management* options should be continuously monitored and reviewed in order to ensure that the objectives are being achieved.

8. **Risk communication**

Communication with all interested parties should be promoted at the earliest opportunity and integrated into all phases of *risk analysis*. This will provide all interested parties, including risk managers, with a better understanding of *risk management* approaches. *Risk communication* should be also well documented.

**Article 6.5.4.**

**Analysis of risks to aquatic animal health**

1. **Definition of the risk**

The *infection* of *aquatic animals* with microorganisms in which resistance has emerged due to antimicrobial usage in *aquatic animals*, and resulting in the loss of benefit of antimicrobial therapy used to manage the *aquatic animal infection*.

2. **Hazard**

- Microorganisms that have acquired resistance (including multiple resistance) arising from the use of an *antimicrobial agent* in *aquatic animals*.
- Microorganisms having obtained a resistance determinant from another microorganism which has acquired resistance arising from the use of an *antimicrobial agent* in *aquatic animals*.

The identification of the hazard should include considerations of the class or subclass of the *antimicrobial agent*. This definition should be read in conjunction with point 3 of Article 6.5.1.

3. **Entry assessment**

The following factors should be considered in the entry assessment:

- *aquatic animal* species treated with the *antimicrobial agent* in question;
- *aquaculture* production system (intensive or extensive, net pens, tanks, raceways, ponds, other);
- number of *aquatic animals* treated, and their age, geographical distribution and, where appropriate, sex;
- prevalence of disease for which the *antimicrobial agent* is indicated or is used in the target *aquatic animal* population;
Chapter 6.5.- Risk analysis for antimicrobial resistance arising from the use of antimicrobial agents in aquatic animals

- data on trends in antimicrobial agent use or sales and changes in aquaculture production systems;
- data on potential extra-label or off-label use;
- methods and routes of administration of the antimicrobial agent;
- dosage regimen (dose, dosing interval and duration of the treatment);
- the pharmacokinetics and pharmacodynamics of the antimicrobial agent;
- type and site of infection;
- development of resistant microorganisms;
- prevalence of pathogenic agents that are likely to develop resistance in an aquatic animal species;
- mechanisms and pathways of direct or indirect transfer of resistance;
- cross-resistance or co-resistance with other antimicrobial agents;
- data on trends and occurrence of resistant microorganisms obtained through surveillance of aquatic animals, aquatic animal products and waste products.

The following confounding factors should be considered in the entry assessment:
- resistant microorganisms or resistant determinants associated with aquatic animals or their products that are a result of terrestrial contamination of the aquatic environment, feed contamination or contamination during post-harvest processing.

4. Exposure assessment

The following factors should be considered in the exposure assessment:
- prevalence and trends of resistant microorganisms in clinically ill and clinically unaffected aquatic animals;
- prevalence of resistant microorganisms in feed and in the aquatic animal environment;
- animal-to-animal transmission of the resistant microorganisms and their resistance determinants (aquatic animal husbandry practices, movement of aquatic animals);
- number or percentage of aquatic animals treated;
- quantity and trends of antimicrobial agent used in aquatic animals;
- survival capacity and spread of resistant microorganisms;
- exposure of wildlife to resistant microorganisms;
- disposal practices for waste products and the likelihood for aquatic animal exposure to resistant microorganisms or resistance determinants through those products;
- capacity of resistant microorganisms to become established in aquatic animals;
- exposure to resistance determinants from other sources such as water, effluent, waste pollution, etc.;
- pharmacokinetics, such as metabolism, bioavailability, distribution to relevant flora - considering the gastrointestinal flora of many aquatic species may be transient;
- transfer of resistant microorganisms and resistance determinants between humans, aquatic animals, and the environment.

5. Consequence assessment

The following factors should be considered in the consequence assessment:
- microbial dose and subsequent host response interactions;
- variation in disease susceptibility of exposed populations and subgroups of the populations;
- variation and frequency of aquatic animal health effects resulting from loss of efficacy of antimicrobial agents and associated costs;
- potential linkage of virulence attributes and resistance;
- importance of the antimicrobial agent in aquatic animal health and human health (see OIE List of Antimicrobial Agents of Veterinary Importance and WHO List of Critically Important Antimicrobials);
- additional burden of disease due to antimicrobial resistant microorganisms;
- number of therapeutic failures due to antimicrobial resistant microorganisms;
- increased severity and duration of infectious disease;
- impact on aquatic animal welfare;
- estimation of the economic impact and cost on aquatic animal health and production;
- deaths (total per year; probability per year for a random member of the population or a member of a specific more exposed sub-population) linked to antimicrobial resistant microorganisms when compared with deaths linked to sensitive microorganisms of the same species;
– availability of alternative antimicrobial therapy;
– potential impact of switching to an alternative antimicrobial agent e.g. alternatives with potential increased toxicity.

6. Risk estimation

A risk estimation integrates the results from the entry assessment, exposure assessment and consequence assessment to produce overall estimates of risks associated with the hazards. Thus, risk estimation takes into account the whole of the risk pathway from hazard identification to the unwanted consequences.

7. Risk management

The relevant provisions in point 7 of Article 6.5.3. apply.

8. Risk communication

The relevant provisions in point 8 of Article 6.5.3. apply.

NB: FIRST ADOPTED IN 2015.
CHAPTER 7.1.

INTRODUCTION TO RECOMMENDATIONS FOR THE WELFARE OF FARMED FISH

Article 7.1.1.

Guiding principles

1) Considering that:
   a) the use of fish in harvest or capture fisheries, in research and for recreation (e.g. ornamentals and aquaria), makes a major contribution to the wellbeing of people; and
   b) there is a critical relationship between fish health and fish welfare; and
   c) improvements in farmed fish welfare can often improve productivity and hence lead to economic benefits.

2) The OIE will develop recommendations for the welfare of farmed fish (excluding ornamental species) during transport, slaughter, and destruction for disease control purposes. In developing these, the following principles will apply:
   a) The use of fish carries with it an ethical responsibility to ensure the welfare of such animals to the greatest extent practicable.
   b) The scientific assessment of fish welfare involves both scientifically derived data and value-based assumptions that need to be considered together, and the process of making these assessments should be made as explicit as possible.

Article 7.1.2.

Scientific basis for recommendations

1) The basic requirements for the welfare of farmed fish include handling methods appropriate to the biological characteristics of the fish and a suitable environment to fulfil their needs.

2) There are many species of fish in farming systems and these have different biological characteristics. It is not practicable to develop specific recommendations for each of these species. These OIE recommendations therefore address the welfare of farmed fish at a general level.
CHAPTER 7.2.

WELFARE OF FARMED FISH DURING TRANSPORT

Article 7.2.1.

Scope

This chapter provides recommendations to minimise the effect of transport on the welfare of farmed fish (hereafter referred to as fish). It applies to their transport by air, by sea or on land within a country and between countries, and only considers the issues related to their welfare.

Recommendations for measures to control the aquatic animal health risks related to the transport of fish are included in Chapter 5.5.

Article 7.2.2.

Responsibilities

All personnel handling fish throughout the transportation process are responsible for ensuring that consideration is given to the potential impact on the welfare of the fish.

1) The responsibilities of the Competent Authority for the exporting and importing jurisdiction include:
   a) establishing minimum standards for fish welfare during transport, including examination before, during and after their transport, appropriate certification, record keeping, awareness and training of personnel involved in transport;
   b) ensuring implementation of the standards, including possible accreditation of transport companies.

2) Owners and managers of fish at the start and at the end of the journey are responsible for:
   a) the general health of the fish and their fitness for transport at the start of the journey and to ensure the overall welfare of the fish during the transport regardless of whether these duties are subcontracted to other parties;
   b) ensuring trained and competent personnel supervise operations at their facilities for fish to be loaded and unloaded in a manner that avoids injury and causes minimum stress;
   c) having a contingency plan available to enable humane killing of the fish at the start and at the end of the journey, as well as during the journey, if required;
   d) ensuring fish have a suitable environment to enter at their destination that ensures their welfare is maintained.

3) Transporters, in cooperation with the farm owner/manager, are responsible for planning the transport to ensure that the transport can be carried out in accordance with fish health and welfare standards including:
   a) using a well maintained vehicle that is appropriate to the species to be transported;
   b) ensuring trained and competent staff are available for loading and unloading, and to ensure swift humane killing of the fish, if required;
   c) having contingency plans to address emergencies and minimise stress during transport;
   d) selecting suitable equipment for loading and unloading of the vehicle.

4) The person in charge of supervising the transport is responsible for all documentation relevant to the transport, and practical implementation of recommendations for welfare of fish during transport.

Article 7.2.3.

Competence

All parties supervising transport activities, including loading and unloading, should have an appropriate knowledge and understanding to ensure that the welfare of the fish is maintained throughout the process. Competence may be gained through formal training and/or practical experience.

1) All persons handling live fish, or who are otherwise responsible for live fish during transport, should be competent in accordance with their responsibilities listed in Article 7.2.2.
2) **Competent Authority**, farm owners/managers, and transport companies have a responsibility in providing training to their respective staff and other personnel.

3) Any necessary training should address species-specific knowledge and may include practical experience on:
   - a) fish behaviour, physiology, general signs of disease and poor welfare;
   - b) operation and maintenance of equipment relevant to fish health and welfare;
   - c) water quality and suitable procedures for water exchange;
   - d) methods of live fish handling during transport, loading and unloading (species-specific aspects when relevant);
   - e) methods for inspection of the fish, management of situations frequently encountered during transport such as changes in water quality parameters, adverse weather conditions, and emergencies;
   - f) methods for the humane killing of fish in accordance with Chapter 7.4.;
   - g) logbooks and record keeping.

**Planning the transport**

1. **General considerations**

   Adequate planning is a key factor affecting the welfare of fish during transportation. The pre-transport preparation, the duration and route of a transport should be determined by the purpose of the transport e.g. biosecurity issues, transport of fish for stocking farms or resource enhancement, for slaughter/killing for disease control purposes. Before the transport starts, plans should be made in relation to:
   - a) type of vehicle and transport equipment required;
   - b) route – such as distance, expected weather and/or sea conditions;
   - c) nature and duration of the transport;
   - d) assessment of the need for acclimatisation of fish to water quality at the site of unloading;
   - e) need for care of the fish during the transport;
   - f) emergency response procedures related to fish welfare;
   - g) assessment of the necessary biosecurity level (e.g. washing and disinfection practices, safe places for changing water, treatment of transport water) (refer to Chapter 5.5.).

2. **Vehicle design and maintenance, including handling equipment**

   - a) **Vehicles and containers** used for transport of fish should be appropriate to the species, size, weight and number of fish to be transported.
   - b) **Vehicles and containers** should be maintained in good mechanical and structural condition to prevent predictable and avoidable damage of the vehicle that may directly or indirectly affect the welfare of transported fish.
   - c) **Vehicles** (if relevant) and **containers** should have adequate circulation of water and equipment for oxygenation as required to meet variations in the conditions during the journey and the needs of the animals being transported, including the closing of valves in well boats for biosecurity reasons.
   - d) The fish should be accessible to inspection en route, if necessary, to ensure that fish welfare can be assessed.
   - e) Documentation that focuses on fish welfare and thus carried with the vehicle should include a transport logbook of stocks received, contact information, mortalities and disposal/storage logs.
   - f) Equipment used to handle fish, for example nets and dip nets, pumping devices and brailing devices, should be designed, constructed and maintained to minimise physical injuries.

3. **Water**

   - a) Water quality (e.g. oxygen, CO₂ and NH₃ level, pH, temperature, salinity) should be appropriate for the species being transported and method of transportation.
   - b) Equipment to monitor and maintain water quality may be required depending on the length of the transport.

4. **Preparation of fish for the transport**

   - a) Prior to transport, feed should be withheld from the fish, taking into consideration the fish species and life stage to be transported.
Chapter 7.2. - Welfare of farmed fish during transport

b) The ability of the fish to cope with the stress of transport should be assessed based on health status, previous handling and recent transport history of the fish. Generally, only fish that are fit for transport should be loaded. Transport for disease control purposes should be in accordance with Chapter 7.4.

c) Reasons for considering of unfitness of fish for transport include:
  
i) displaying clinical signs of disease;
  
ii) significant physical injuries or abnormal behaviour, such as rapid ventilation or abnormal swimming;
  
iii) recent exposure to stressors that adversely affect behaviour or physiological state (for example extreme temperatures, chemical agents);
  
iv) insufficient or excessive length of fasting.

5. Species-specific recommendations

Transport procedures should take account of variations in the behaviour and specific needs of the transported fish species. Handling procedures that are successful with one species may be ineffective or dangerous for another species.

Some species or life stages may need to be physiologically prepared prior to entering a new environment, such as by feed deprivation or osmotic acclimatisation.

6. Contingency plans

There should be a contingency plan that identifies the important adverse fish welfare events that may be encountered during the transport, the procedures for managing each event and the action to be taken in such an event. For each event, the plan should document the actions to be undertaken and the responsibilities of all parties involved, including communications and record keeping.

Article 7.2.5.

Documentation

1) Fish should not be loaded until the required documentation is complete.

2) The documentation accompanying the consignment (the transport log) should include:
   
a) description of the consignment (e.g. date, time, and place of loading, species, biomass load);
   
b) description of the transport plan (e.g. including route, water exchanges, expected time, date and place of arrival and unloading and receiver contact information).

3) The transport log should be made available to the dispatcher and the receiver of the consignment as well as to the Aquatic Animal Health Service upon request. Transport logs from previous journeys should be kept after completion of the transport for a period of time as specified by the Aquatic Animal Health Service.

Article 7.2.6.

Loading the fish

1) The issues which should be addressed to avoid injury and unnecessary stress to the fish include:
   
a) crowding procedure in farm pond, tank, net or cage prior to loading;
   
b) equipment (such as nets, pumps, pipes and fittings) that are improperly constructed (e.g. sharp bends or protrusions) or improperly operated (e.g. overloading with fish of incorrect size or number of fish);
   
c) water quality - some species of fish should be acclimatised if there is a likelihood of the fish being transported in water of a significantly different temperature or other water parameters.

2) The density of fish in a vehicle and/or container should be in accordance with scientific data where available and not exceed what is generally accepted for a given species and a given situation.

3) Loading should be carried out, or supervised, by operators with knowledge and experience of the behaviour and other characteristics of the fish species being loaded to ensure that the welfare of the fish is maintained.
Article 7.2.7.

Transporting the fish

1. General considerations
   a) Periodic inspections should take place during the transport to verify that acceptable welfare is being maintained.
   b) Ensure that water quality is monitored and the necessary adjustments made to avoid extreme conditions.
   c) Travel in a manner that minimises uncontrolled movements of the fish that may lead to stress and cause injury.

2. Sick or injured fish
   a) In the event of a fish health emergency during transport, the vehicle operator should initiate the contingency plan (see point 6 of Article 7.2.4.).
   b) If the killing of fish is necessary during the transport, it should be carried out humanely in accordance with Chapter 7.4.

Article 7.2.8.

Unloading the fish

1) The principles of good fish handling during loading apply equally during unloading.
2) Fish should be unloaded as soon as possible after arrival at the destination, allowing sufficient time to ensure that the unloading procedure does not cause harm to the fish. Some species of fish should be acclimatised if there is a likelihood of the fish being unloaded into water of a significantly different quality (such as temperature, salinity, pH).
3) Moribund or seriously injured fish should be removed and humanely killed in accordance with Chapter 7.4.

Article 7.2.9.

Post-transport activities

1) The person in charge of receiving the fish should closely observe them during the post-transport period, and keep appropriate records.
2) Fish showing abnormal clinical signs should be humanely killed in accordance with Chapter 7.4 or isolated and examined by a veterinarian or other qualified personnel, who may recommend treatment.
3) Significant problems associated with transport should be evaluated to prevent recurrence of such problems.

NB: FIRST ADOPTED IN 2009; MOST RECENT UPDATE ADOPTED IN 2012.
CHAPTER 7.3.

WELFARE ASPECTS OF STUNNING AND KILLING OF FARMED FISH FOR HUMAN CONSUMPTION

Article 7.3.1.

Scope

These recommendations apply to the stunning and killing of farmed fish species for human consumption.

These recommendations address the need to ensure the welfare of farmed fish, intended for human consumption, during stunning and killing including transport and holding immediately prior to stunning.

This chapter describes general principles that should be applied to ensure the welfare of fish for stunning and killing for human consumption and also applies to farmed fish killed for disease control purposes. Other measures applicable to emergency killing for disease control purposes are addressed in Chapter 7.4.

As a general principle, farmed fish should be stunned before killing, and the stunning method should ensure immediate and irreversible loss of consciousness. If the stunning is not irreversible, fish should be killed before consciousness is recovered.

Article 7.3.2.

Personnel

Persons engaged in the handling, stunning and killing of fish play an important role in their welfare. Personnel handling fish for stunning and killing should be experienced and competent in the handling of fish, and understand their behaviour patterns as well as the underlying principles necessary to carry out their tasks. Some stunning and killing methods may pose a risk to the personnel; therefore training should cover occupational health and safety implications of any methods used.

Article 7.3.3.

Transport

If fish are to be transported prior to stunning and killing, this should be done in accordance with OIE recommendations on the welfare of farmed fish during transport (see Chapter 7.2.).

Article 7.3.4.

Design of holding facilities

1) The holding facilities should be designed and specifically constructed to hold a certain fish species or group of fish species.

2) The holding facilities should be of a size that allows holding a certain number of fish for processing in a given timeframe without compromising the welfare of the fish.

3) Operations should be conducted with minimal injury and stress to the fish.

4) The following recommendations may help to achieve this:
   a) nets and tanks should be designed and maintained to minimise physical injuries;
   b) water quality should be suitable for the fish species and stocking density;
   c) equipment for transferring fish, including pumps and pipes, should be designed and maintained to minimise injury.
Chapter 7.3.- Welfare aspects of stunning and killing of farmed fish for human consumption

Article 7.3.5.

Unloading, transferring and loading

1) Fish should be unloaded, transferred and loaded under conditions that minimise injury and stress to the fish.

2) The following points should be considered:
   a) Water quality (e.g. temperature, oxygen and $\text{CO}_2$ levels, pH and salinity) should be assessed on arrival of fish prior to their unloading, and corrective action taken if required.
   b) Where possible any injured or moribund fish should be separated and killed humanely.
   c) The crowding periods of fish should be as short and infrequent as possible to avoid stressful conditions arising.
   d) The handling of fish during transfers should be minimised and preferably fish should not be handled out of water. If fish need to be removed from water, this period should be kept as short as possible.
   e) Where feasible, and when applicable, fish should be allowed to swim directly into a stunning device without handling to avoid handling stress.
   f) Equipment used to handle fish, for example nets and dip nets, pumping devices and brailing devices, should be designed, constructed and operated to minimise physical injuries (e.g. pumping height, pressure and speed are important factors to consider).
   g) Fish should not be fasted (deprived of food) before killing for longer than is necessary, e.g. to clear the gut or to reduce undesirable organoleptic properties.
   h) There should be a contingency plan to address emergencies and minimise stress during unloading, transferring and loading fish.

Article 7.3.6.

Stunning and killing methods

1. General considerations

   a) The choice of method should take account of species-specific information where available.
   b) All handling, stunning and killing equipment should be maintained and operated appropriately; it should be tested on a regular basis to ensure that performance is adequate.
   c) Effective stunning should be verified by the absence of consciousness.
   d) A backup stunning system is necessary. Any fish mis-stunned, or regaining consciousness before death, should be re-stunned as soon as possible.
   e) Stunning should not take place if killing is likely to be delayed such that the fish will recover or partially recover consciousness.
   f) While absence of consciousness may be difficult to recognise, signs of correct stunning include i) loss of body and respiratory movement (loss in opercular activity); ii) loss of visual evoked response (VER); iii) loss of vestibulo-ocular reflex (VOR, eye rolling).

2. Mechanical stunning and killing methods

   a) Percussive stunning is achieved by a blow of sufficient strength to the head applied above or immediately adjacent to the brain in order to damage the brain. Mechanical stunning may be achieved either manually or using specially developed equipment.
   b) Spiking or coring are irreversible stunning and killing methods of fish based on physical damage to the brain by inserting a spike or core into the brain.
   c) Shooting using a free bullet may be used for killing large fish (such as tuna). The fish may either be crowded in a net and shot in the head from the surface, or individual fish may be killed by shooting in the head from under the water (commonly called lupara).
   d) Unconsciousness following mechanical stunning is generally irreversible if correctly applied. In cases where the loss of consciousness is transient, fish should be killed before consciousness is recovered.
3. Electrical stunning and killing methods
   a) Electrical stunning involves the application of an electrical current of sufficient strength and duration, and suitable frequency to cause immediate loss of consciousness and insensibility of the fish. The conductivity of fresh and brackish water varies, so it is essential to establish the parameters of the electrical current to ensure proper stunning at the site of stunning.

   b) The electrical stunning device should be constructed and used for the specific fish species and their environment.

   c) Unconsciousness following electrical stunning may be reversible. In such cases fish should be killed before consciousness is recovered.

   d) Fish should be confined beneath the surface of the water, and there should be a uniform distribution of electrical current in the stunning tank or chamber.

   e) In semi-dry electrical stunning systems, fish should enter the device head first to ensure rapid and efficient stunning.

4. Other killing methods
   The following methods are known to be used for killing fish: chilling with ice in holding water, carbon dioxide (CO₂) in holding water; chilling with ice and CO₂ in holding water; salt or ammonia baths; asphyxiation by removal from water; exsanguination without stunning. However, they have been shown to result in poor fish welfare. Therefore, these methods should not be used if it is feasible to use the methods described in points 2 and 3 of this article, as appropriate to the fish species.

Article 7.3.7.

Summary table of some stunning/killing methods for fish and their respective welfare issues

<table>
<thead>
<tr>
<th>Stunning/killing method</th>
<th>Specific method</th>
<th>Key fish welfare concerns/requirements</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical</td>
<td>Spiking or coring</td>
<td>The spike should be aimed on the skull in a position to penetrate the brain of the fish and the impact of the spike should produce immediate unconsciousness. Fish should be quickly removed from the water, restrained and the spike immediately inserted into the brain. It is a stun / kill method.</td>
<td>Immediate loss of consciousness. Suitable for medium to large sized fish.</td>
<td>Inaccurate application may cause injuries. Difficult to apply if fish agitated. It is only practicable for the killing of a limited number of fish.</td>
</tr>
<tr>
<td></td>
<td>Free bullet</td>
<td>The shot should be carefully aimed at the brain. The fish should be positioned correctly and the shooting range should be as short as practicable. It is a stun / kill method.</td>
<td>Immediate loss of consciousness. Suitable for large sized fish (e.g. large tuna).</td>
<td>Shooting distance; calibre need to be adapted. Excessive crowding and noise of guns may cause stress reaction. Contamination of the working area due to release of body fluids may present a biosecurity risk. May be hazardous to operators.</td>
</tr>
<tr>
<td>Mechanical</td>
<td>Percussive stunning</td>
<td>The blow should be of sufficient force and delivered above or adjacent to the brain in order to render immediate unconsciousness. Fish should be quickly removed from the water, restrained and given a quick blow to the head, delivered either manually by a club or by automated percussive stunning. The effectiveness of stunning should be checked, and fish be re-stunned if necessary. It can be a stun / kill method.</td>
<td>Immediate loss of consciousness. Suitable for medium to large sized fish.</td>
<td>Hand operated equipment may be hampered by uncontrolled movement of the fish. Mis-stunning may result from a too weak blow. Injuries may occur. Manual percussive stunning is only practicable for the killing of a limited number of fish of a similar size.</td>
</tr>
</tbody>
</table>
Chapter 7.3.- Welfare aspects of stunning and killing of farmed fish for human consumption

<table>
<thead>
<tr>
<th>Stunning/killing method</th>
<th>Specific method</th>
<th>Key fish welfare concerns/requirements</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrical</td>
<td>Electrical stunning</td>
<td>Involves the application of an electrical current of sufficient strength, frequency and duration to cause immediately unconsciousness. It can be a stun / kill method. Equipment should be designed and maintained correctly.</td>
<td>Immediate loss of consciousness. Suitable for small to medium sized fish. Suitable for large numbers of fish, and the fish do not have to be removed from the water.</td>
<td>Difficult to standardise for all species. Optimal control parameters are unknown for some species. May be hazardous to operators.</td>
</tr>
<tr>
<td></td>
<td>Semi-dry electrical stunning</td>
<td>The head of the fish should enter the system first so electricity is applied to the brain first. Involves the application of an electrical current of sufficient strength, frequency and duration to cause immediately unconsciousness. Equipment should be designed and maintained correctly.</td>
<td>Good visual control of stunning and the ability for re-stunning of individual fish.</td>
<td>Misplacement of the fish may result in improper stunning. Optimal control parameters are unknown for some species. Not suitable for mixed sizes of fish.</td>
</tr>
</tbody>
</table>

[Note: the terms small, medium and large fish should be interpreted relative to the species in question.]

Article 7.3.8.

Examples of stunning/killing methods for fish groups

The following methods enable humane killing for the following fish groups:

1) percussive stunning: carp, salmonids;
2) spiking or coring: tuna;
3) free bullet: tuna;
4) electrical stunning: carp, eel, salmonids.

NB: FIRST ADOPTED IN 2010; MOST RECENT UPDATE ADOPTED IN 2012.
CHAPTER 7.4.

KILLING OF FARMED FISH FOR DISEASE CONTROL PURPOSES

Article 7.4.1.

Scope

These recommendations are based on the premise that a decision to kill the farmed fish for disease control purposes has been made, and address the need to ensure the welfare of the farmed fish until they are dead.

The culling of individual farmed fish, in the course of farming operations (i.e. sorting, grading, or background morbidity), is out of the scope of this chapter.

Account should also be taken of the guidance given in the following chapters in the Aquatic Code: 4.5. Contingency planning; Chapter 4.7. Handling, disposal and treatment of aquatic animal waste; Chapter 5.5. Control of aquatic animal health risks associated with transport; Chapter 7.2. Welfare of farmed fish during transport and Chapter 7.3. Welfare aspects of stunning and killing of farmed fish for human consumption.

Article 7.4.2.

General principles

1) Fish welfare considerations should be addressed within contingency plans for disease control (refer to Chapter 4.5.).

2) The killing method should be selected taking into consideration fish welfare and biosecurity requirements as well as safety of the personnel.

3) When fish are killed for disease control purposes, methods used should result in immediate death or immediate loss of consciousness lasting until death; when loss of consciousness is not immediate, induction of unconsciousness should be non-aversive or the least aversive possible and should not cause avoidable pain, distress or suffering in fish.

4) The methods described in Chapter 7.3. can also be used for disease control purposes.

5) Some of the methods recommended for disease control purposes (e.g. anaesthetic overdose, maceration) may render the fish unsuitable for human consumption, and this should be specified in the contingency plan.

6) Depending on the situation, emergency killing of fish may be carried out on site or after fish are transported to an approved killing facility.

Article 7.4.3.

Operational guidelines for affected premises and approved killing facilities

1) The following should apply when killing fish:
   a) Operational procedures should be adapted to the specific circumstances on the premises and should address fish welfare and biosecurity specific to the disease of concern.
   b) Killing of fish should be carried out without delay by appropriately qualified personnel with all due consideration made to increased biosecurity protocols.
   c) Handling of fish should be kept to a minimum to avoid stress and to prevent spread of disease. This should be done in accordance with the articles described below.
   d) Methods used to kill the fish should render them unconscious until death or kill them in the shortest time possible, and should not cause avoidable pain or distress.
   e) There should be continuous monitoring of the procedures to ensure they are consistently effective with regard to biosecurity and fish welfare.
   f) Standard operating procedures (SOP's) should be available and followed at the premises.
Chapter 7.4.- Killing of farmed fish for disease control purposes

2) Procedures for the killing of fish on affected premises for disease control purposes should be developed by the operator and approved by the Competent Authority, taking into consideration fish welfare and biosecurity requirements as well as safety of the personnel and should include consideration of:

a) handling and movement of fish;

b) species, number, age and size of fish to be killed;

c) methods for killing the fish;

d) availability of anaesthetic agents suitable to kill the fish;

e) equipment needed to kill the fish;

f) any legal issues (e.g. the use of anaesthetic agents suitable for killing fish);

g) presence of other nearby aquaculture premises;

h) disposal of killed fish in accordance with Chapter 4.7.

Article 7.4.4.

Competencies and responsibilities of the operational team

The operational team is responsible for planning, implementation of, and reporting on the killing of the fish.

1. Team leader

a) Competencies

i) Ability to assess fish welfare, especially relating to the effectiveness of the stunning and killing techniques selected and utilised in the fish killing operations, to detect and correct any deficiencies;

ii) ability to assess biosecurity risks and mitigation measures being applied to prevent spread of disease;

iii) skills to manage all activities on premises and deliver outcomes on time;

iv) awareness of the psychological impact on fish farmers, team members and general public;

v) effective communication skills.

b) Responsibilities

i) Determine most appropriate killing method(s) to ensure that the fish are killed without avoidable pain and distress while balancing biosecurity considerations;

ii) plan overall operations on the affected premises;

iii) determine and address requirements for fish welfare, operator safety and biosecurity;

iv) organise, brief and manage a team of people to facilitate killing of the relevant fish in accordance with national contingency plans for disease control;

v) determine logistics required;

vi) monitor operations to ensure that fish welfare, operator safety and biosecurity requirements are met;

vii) report upwards on progress and problems;

viii) provide a written report summarising the killing practices utilised in the operation and their effect on fish welfare and subsequent biosecurity outcomes. The report should be archived and be accessible for a period of time defined by the Competent Authority;

ix) review on-site facilities in terms of their appropriateness for mass destruction.

2. On-site personnel responsible for killing of fish

a) Competencies

i) Specific knowledge of fish, their behaviour and environment;

ii) trained and competent in fish handling, stunning and killing procedures;

iii) trained and competent in the operation and maintenance of equipment.

b) Responsibilities

i) Ensure killing of fish through effective stunning and killing techniques;

ii) assist team leader as required;

iii) design and construct temporary fish handling facilities, when required.
Chapter 7.4.- Killing of farmed fish for disease control purposes

Article 7.4.5.

Killing by an overdose of an anaesthetic agent

This article refers to killing methods using an overdose of an anaesthetic agent.

1. Use of anaesthetic agents
   a) Anaesthetic agents used for killing fish should kill the fish effectively, not merely have an anaesthetic effect.
   b) When using anaesthetic agents, the operating personnel should ensure that the solution has the correct concentration for the water in which it is to be administered, and that water of appropriate quality for the species and life stage of fish is used.
   c) Fish should be kept in the anaesthetic solution until they are dead.

2. Advantages
   a) Large numbers of fish may be killed in one batch.
   b) Handling is not required until fish are dead.
   c) Use of anaesthetic agents is a non-invasive technique and thus reduces biosecurity risks.

3. Disadvantages
   a) The method may fail to cause death in fish, e.g. dilution of the anaesthetic solution with prolonged use. In such circumstances, fish that are anaesthetised should be killed before they regain consciousness.
   b) Some anaesthetic agents may induce a transient aversive reaction in the fish.
   c) Care is essential in the preparation and provision of treated water, and in the disposal of water and/or fish carcasses that have been treated with anaesthetic agents.

Article 7.4.6.

Mechanical killing methods

1. Decapitation
   a) Decapitation, using a sharp device, such as a guillotine or knife, may be used but should be preceded by stunning or, if appropriate, anaesthesia.
   b) The required equipment should be kept in good working order.
   c) Contamination of the working area by blood, body fluids and other organic material may present a biosecurity risk and is the major disadvantage of this method.

2. Maceration
   a) Maceration by a mechanical device with rotating blades or projections causes immediate fragmentation and death in newly hatched fish and embryonated eggs, as well as fertilised/unfertilised eggs of fish. It is a suitable method for the processing of such material. A large number of eggs/newly hatched fry can be killed quickly.
   b) Maceration requires specialised equipment which should be kept in good working order. The rate of introducing material into the device should be such that the cutting blades continue to rotate at their fully functional rate and that they do not fall below the defined critical speed defined by the manufacturer.
   c) Contamination of the working area by blood, body fluids and other organic material may present a biosecurity risk and is the major disadvantage of this method.

NB: FIRST ADOPTED IN 2012; MOST RECENT UPDATE ADOPTED IN 2013.
SECTION 8.
DISEASES OF AMPHIBIANS

CHAPTER 8.1.
INFECTION WITH BATRACHOCHYTRIUM DENDROBATIDIS

Article 8.1.1.

For the purposes of the Aquatic Code, infection with Batrachochytrium dendrobatidis means infection with the freshwater fungus *B. dendrobatidis* (Fungi, Chytridiomycota, Rhizophydiales).

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 8.1.2.

Scope

The recommendations in this chapter apply to: all species of *Anura* (frogs and toads), *Caudata* (salamanders, newts and sirens) and *Gymnophiona* (caecilians). The recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 8.1.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with *B. dendrobatidis* status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with *B. dendrobatidis*, regardless of the infection with *B. dendrobatidis* status of the exporting country, zone or compartment, when authorising the importation or transit of the following *aquatic animal products* from the species referred to in Article 8.1.2. which are intended for any purpose and which comply with Article 5.4.1.:
   a) heat sterilised hermetically sealed amphibian products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) cooked amphibian products that have been subjected to heat treatment at 100°C for at least one minute (or any time/temperature equivalent which has been demonstrated to inactivate *B. dendrobatidis*);
   c) pasteurised amphibian products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent which has been demonstrated to inactivate *B. dendrobatidis*);
   d) mechanically dried amphibian products (i.e. a heat treatment of 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate *B. dendrobatidis*);
   e) amphibian skin leather.

2) When authorising the importation or transit of *aquatic animals* and *aquatic animal products* of a species referred to in Article 8.1.2., other than those referred to in point 1 of Article 8.1.3., Competent Authorities should require the conditions prescribed in Articles 8.1.7. to 8.1.12. relevant to the infection with *B. dendrobatidis* status of the exporting country, zone or compartment.
3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 8.1.2. but which could reasonably be expected to pose a risk of spread of infection with *B. dendrobatidis*, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

**Article 8.1.4.**

**Country free from infection with *B. dendrobatidis***

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with *B. dendrobatidis* if all the areas covered by the zone are declared infection with *B. dendrobatidis* free (see Article 8.1.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with *B. dendrobatidis* if:

1) none of the susceptible species referred to in Article 8.1.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 8.1.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with *B. dendrobatidis*;

OR

4) it previously made a self-declaration of freedom from infection with *B. dendrobatidis* and subsequently lost its disease free status due to the detection of infection with *B. dendrobatidis* but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with *B. dendrobatidis*.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 8.1.5.

**Article 8.1.5.**

**Zone or compartment free from infection with *B. dendrobatidis***

If a zone or compartment extends over more than one country, it can only be declared an infection with *B. dendrobatidis* free zone or compartment if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with *B. dendrobatidis* may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 8.1.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;
Chapter 8.1.- Infection with Batrachochytrium dendrobatidis

OR
2) any of the susceptible species referred to in Article 8.1.2. are present in the zone or compartment and the following conditions have been met;
   a) there has not been any observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR
3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with B. dendrobatidis;

OR
4) it previously made a self-declaration of freedom from infection with B. dendrobatidis and subsequently lost its disease free status due to the detection of infection with B. dendrobatidis but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with B. dendrobatidis.

Article 8.1.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with B. dendrobatidis following the provisions of points 1 or 2 of Articles 8.1.4. or 8.1.5. (as relevant) may maintain its status as free from infection with B. dendrobatidis provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with B. dendrobatidis following the provisions of point 3 of Articles 8.1.4. or 8.1.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with B. dendrobatidis, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with B. dendrobatidis, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 8.1.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with B. dendrobatidis

When importing aquatic animals and aquatic animal products of species referred to in Article 8.1.2. from a country, zone or compartment declared free from infection with B. dendrobatidis, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 8.1.4. or 8.1.5. (as applicable) and 8.1.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with B. dendrobatidis.
Chapter 8.1.- Infection with Batrachochytrium dendrobatidis

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 8.1.3.

Article 8.1.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with B. dendrobatidis

When importing for aquaculture, aquatic animals of a species referred to in Article 8.1.2. from a country, zone or compartment not declared free from infection with B. dendrobatidis, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive B. dendrobatidis in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with B. dendrobatidis.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for B. dendrobatidis in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with B. dendrobatidis (as described in Chapter 2.1.1. of the Aquatic Manual) and test for B. dendrobatidis in accordance with Chapter 1.4.;
      v) if B. dendrobatidis is not detected in the F-1 population, it may be defined as free from infection with B. dendrobatidis and may be released from quarantine;
      vi) if B. dendrobatidis is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 8.1.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with B. dendrobatidis

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 8.1.2. from a country, zone or compartment not declared free from infection with B. dendrobatidis, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 8.1.3., or products described in point 1 of Article 8.1.12., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of B. dendrobatidis or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 8.1.- Infection with Batrachochytrium dendrobatidis

Article 8.1.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with B. dendrobatidis

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 8.1.2. from a country, zone or compartment not declared free from infection with B. dendrobatidis, the Competent Authority of the importing country should require that:

1) the consignment be delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water and equipment used in transport and all effluent and waste materials from the processing facility be treated in a manner that inactivates B. dendrobatidis.

This article does not apply to commodities referred to in point 1 of Article 8.1.3.

Article 8.1.11.

Importation of aquatic animals intended for use in laboratories or zoos from a country, zone or compartment not declared free from infection with B. dendrobatidis

When importing, for use in laboratories or zoos, aquatic animals of species referred to in Article 8.1.2. from a country, zone or compartment not declared free from infection with B. dendrobatidis, the Competent Authority of the importing country should ensure:

1) the direct delivery to and lifelong holding of the consignment in quarantine facilities authorised by the Competent Authority; and

2) the treatment of water and equipment used in transport and of all effluent and waste materials in a manner that inactivates B. dendrobatidis; and

3) the disposal of carcasses in accordance with Chapter 4.7.

Article 8.1.12.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with B. dendrobatidis

1) Competent Authorities should not require any conditions related to infection with B. dendrobatidis, regardless of the infection with B. dendrobatidis status of the exporting country, zone or compartment, when authorising the importation or transit of amphibian meat (skin off and fresh or frozen) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 8.1.2. from a country, zone or compartment not declared free from infection with B. dendrobatidis, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2008; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 8.2.

INFECTION WITH RANAVIRUS

Article 8.2.1.

For the purposes of the Aquatic Code, infection with ranavirus means infection with any member virus species of the genus Ranavirus in the family Iridoviridae with the exception of epizootic haematopoietic necrosis virus and European catfish virus.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 8.2.2.

Scope

The recommendations in this chapter apply to: all species of Anura (frogs and toads) and Caudata (salamanders and newts). The recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 8.2.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with ranavirus status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with ranavirus, regardless of the infection with ranavirus status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 8.2.2. which are intended for any purpose and which comply with Article 5.4.1.:

   a) heat sterilised hermetically sealed amphibian products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   
   b) cooked amphibian products that have been subjected to heat treatment at 65°C for at least 30 minutes (or any time/temperature equivalent which has been demonstrated to inactivate all virus species of the genus Ranavirus in the family Iridoviridae [with the exception of epizootic haematopoietic necrosis virus and European catfish virus]);
   
   c) pasteurised amphibian products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent which has been demonstrated to inactivate all virus species of the genus Ranavirus in the family Iridoviridae [with the exception of epizootic haematopoietic necrosis virus and European catfish virus]);
   
   d) mechanically dried amphibian products (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate all virus species of the genus Ranavirus in the family Iridoviridae [with the exception of epizootic haematopoietic necrosis virus and European catfish virus]).

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 8.2.2., other than those referred to in point 1 of Article 8.2.3., Competent Authorities should require the conditions prescribed in Articles 8.2.7. to 8.2.12. relevant to the infection with ranavirus status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 8.2.2. but which could reasonably be expected to pose a risk of spread of infection with ranavirus, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.
Article 8.2.4.

**Country free from infection with ranavirus**

If a country shares a *zone* with one or more other countries, it can only make a *self-declaration of freedom* from infection with ranavirus if all the areas covered by the *zone* are declared free from infection with ranavirus (see Article 8.2.5.).

As described in Article 1.4.6., a country may make a *self-declaration of freedom* from infection with ranavirus if:

1) none of the *susceptible species* referred to in Article 8.2.2. are present and *basic biosecurity conditions* have been continuously met for at least the last two years;

OR

2) any of the *susceptible species* referred to in Article 8.2.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the *disease* for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the *Aquatic Manual*);
   and
   b) *basic biosecurity conditions* have been continuously met for at least the last ten years;

OR

3) the disease status prior to *targeted surveillance* is unknown but the following conditions have been met:
   a) *basic biosecurity conditions* have been continuously met for at least the last two years; and
   b) *targeted surveillance*, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with ranavirus;

OR

4) it previously made a *self-declaration of freedom* from infection with ranavirus and subsequently lost its *disease* free status due to the detection of infection with ranavirus but the following conditions have been met:
   a) on detection of the *disease*, the affected area was declared an *infected zone* and a *protection zone* was established; and
   b) infected populations have been destroyed or removed from the *infected zone* by means that minimise the *risk* of further spread of the *disease*, and the appropriate *disinfection* procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing *basic biosecurity conditions* have been reviewed and modified as necessary and have continuously been in place since eradication of the *disease*; and
   d) *targeted surveillance*, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with ranavirus.

In the meantime, part or all of the non-affected area may be declared a free *zone* provided that such a part meets the conditions in point 3 of Article 8.2.5.

Article 8.2.5.

**Zone or compartment free from infection with ranavirus**

If a *zone* or *compartment* extends over more than one country, it can only be declared an infection with ranavirus free *zone* or *compartment* if all the relevant *Competent Authorities* confirm that all relevant conditions have been met.

As described in Article 1.4.6., a *zone* or *compartment* within the *territory* of one or more countries not declared free from infection with ranavirus may be declared free by the *Competent Authority(ies)* of the country(ies) concerned if:

1) none of the *susceptible species* referred to in Article 8.2.2. are present in the *zone* or *compartment* and *basic biosecurity conditions* have been continuously met for at least the last two years;

OR

2) any of the *susceptible species* referred to in Article 8.2.2. are present in the *zone* or *compartment* and the following conditions have been met:
   a) there has not been any observed occurrence of the *disease* for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the *Aquatic Manual*); and
   b) *basic biosecurity conditions* have been continuously met for at least the last ten years;
Chapter 8.2.- Infection with ranavirus

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with ranavirus;

OR

4) it previously made a self-declaration of freedom for a zone from infection with ranavirus and subsequently lost its disease free status due to the detection of the infection with ranavirus in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with ranavirus.

Article 8.2.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with ranavirus following the provisions of points 1 or 2 of Articles 8.2.4. or 8.2.5. (as relevant) may maintain its status as free from infection with ranavirus provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with ranavirus following the provisions of point 3 of Articles 8.2.4. or 8.2.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with ranavirus provided that conditions that are conducive to clinical expression of infection with ranavirus, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with ranavirus, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 8.2.7.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with ranavirus

When importing aquatic animals and aquatic animal products of species referred to in Article 8.2.2. from a country, zone or compartment declared free from infection with ranavirus, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 8.2.4. or 8.2.5. (as applicable) and 8.2.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with ranavirus.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 8.2.3.
Chapter 8.2.- Infection with ranavirus

Article 8.2.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with ranavirus

When importing for aquaculture, aquatic animals of a species referred to in Article 8.2.2. from a country, zone or compartment not declared free from infection with ranavirus, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive ranavirus in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with ranavirus.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for ranavirus in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with ranavirus (as described in Chapter 2.1.2. of the Aquatic Manual) and test for ranavirus in accordance with Chapter 1.4.;
      v) if ranavirus is not detected in the F-1 population, it may be defined as free from infection with ranavirus and may be released from quarantine;
      vi) if ranavirus is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 8.2.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with ranavirus

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 8.2.2. from a country, zone or compartment not declared free from infection with ranavirus, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 8.2.3., or products described in point 1 of Article 8.2.12., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of ranavirus or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Article 8.2.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with ranavirus

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of the species referred to in Article 8.2.2. from a country, zone or compartment not declared free from infection with ranavirus, the Competent Authority of the importing country should require that:

1) the consignment be delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and
2) water and equipment used in transport and all effluent and waste materials from the processing facility be treated in a manner that inactivates ranavirus.

This article does not apply to commodities referred to in point 1 of Article 8.2.3.

Article 8.2.11.

Importation of aquatic animals intended for use in laboratories or zoos, from a country, zone or compartment not declared free from infection with ranavirus

When importing, for use in laboratories or zoos, aquatic animals of species referred to in Article 8.2.2. from a country, zone or compartment not declared free from infection with ranavirus, the Competent Authority of the importing country should ensure:

1) the direct delivery to and lifelong holding of the consignment in quarantine facilities authorised by the Competent Authority; and
2) the treatment of water and equipment used in transport and of all effluent and waste materials in a manner that inactivates ranavirus; and
3) the disposal of carcasses in accordance with Chapter 4.7.

Article 8.2.12.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with ranavirus

1) Competent Authorities should not require any conditions related to infection with ranavirus, regardless of the infection with ranavirus status of the exporting country, zone or compartment, when authorising the importation or transit of the following commodities which have been prepared and packaged for retail trade and which comply with Article 5.4.2.:
   – no commodities listed.
2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 8.2.2. from a country, zone or compartment not declared free from infection with ranavirus, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2008; MOST RECENT UPDATE ADOPTED IN 2017.
SECTION 9.
DISEASES OF CRUSTACEANS

CHAPTER 9.1.
ACUTE HEPATOPANCREATIC NECROSIS DISEASE

Article 9.1.1.

For the purposes of the Aquatic Code, acute hepatopancreatic necrosis disease (AHPND) means infection with strains of *Vibrio parahaemolyticus* (Vp AHPND) of the Family Vibrionaceae, that contain a ~70-kbp plasmid with genes that encode homologues of the *Photorhabdus* insect-related (Pir) toxins, PirA and PirB.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 9.1.2.

Scope

The recommendations in this chapter apply to the following species that meet the criteria for listing as susceptible in accordance with Chapter 1.5.: giant tiger prawn (*Penaeus monodon*) and whiteleg shrimp (*Penaeus vannamei*).

Article 9.1.3.

Importation or transit of aquatic animal products for any purpose regardless of the AHPND status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to AHPND, regardless of the AHPND status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.1.2., which are intended for any purpose and comply with Article 5.4.1.:  
   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate Vp AHPND);  
   b) cooked crustacean products that have been subjected to heat treatment at 100°C for at least one minute (or any time/temperature equivalent that has been demonstrated to inactivate Vp AHPND);  
   c) crustacean oil;  
   d) crustacean meal;  
   e) chemically extracted chitin.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.1.2., other than those referred to in point 1 of Article 9.1.3., Competent Authorities should require the conditions prescribed in Articles 9.1.7. to 9.1.11. relevant to the AHPND status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.1.2. but which could reasonably be expected to pose a risk of transmission of AHPND, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.
Chapter 9.1.- Acute hepatopancreatic necrosis disease

Article 9.1.4.

Country free from AHPND

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from AHPND if all the areas covered by the shared water bodies are declared countries or zones free from AHPND (see Article 9.1.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from AHPND if:

1) none of the susceptible species referred to in Article 9.1.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.1.2. are present and the following conditions have been met:
   a) there has been no occurrence of AHPND for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the AHPND status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of AHPND;

OR

4) it previously made a self-declaration of freedom from AHPND and subsequently lost its free status due to the detection of AHPND but the following conditions have been met:
   a) on detection of AHPND, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of AHPND, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of AHPND; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of AHPND.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.1.5.

Article 9.1.5.

Zone or compartment free from AHPND

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from AHPND if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from AHPND may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.1.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.1.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of AHPND for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;
OR
3) the AHPND status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of AHPND;

OR
4) it previously made a self-declaration of freedom for a zone from AHPND and subsequently lost its free status due to the detection of AHPND in the zone but the following conditions have been met:
   a) on detection of AHPND, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of AHPND, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of AHPND; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of AHPND.

Article 9.1.6.

Maintenance of free status

A country, zone or compartment that is declared free from AHPND following the provisions of points 1 or 2 of Articles 9.1.4. or 9.1.5. (as relevant) may maintain its status as free from AHPND provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from AHPND following the provisions of point 3 of Articles 9.1.4. or 9.1.5. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions are conducive to clinical expression of AHPND, as described in the corresponding chapter of the Aquatic Manual, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of AHPND, targeted surveillance should be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 9.1.7.

Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from AHPND

When importing aquatic animals of a species referred to in Article 9.1.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from AHPND, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country. The international aquatic animal health certificate should state that, on the basis of the procedures described in Articles 9.1.4. or 9.1.5. (as applicable) and 9.1.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from AHPND.

The international aquatic animal health certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.1.3.
Chapter 9.1.- Acute hepatopancreatic necrosis disease

Article 9.1.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from AHPND

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.1.2. from a country, zone or compartment not declared free from AHPND, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate VP<sub>AHPND</sub> in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for AHPND.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for VP<sub>AHPND</sub> in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of AHPND (as described in Chapter 2.2.1. of the Aquatic Manual) and test for VP<sub>AHPND</sub> in accordance with Chapter 1.4.;
      v) if VP<sub>AHPND</sub> is not detected in the F-1 population, it may be defined as free from AHPND and may be released from quarantine;
      vi) if VP<sub>AHPND</sub> is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.1.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from AHPND

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.1.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from AHPND, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.1.3. or in point 1 of Article 9.1.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of VP<sub>AHPND</sub> or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of VP<sub>AHPND</sub> or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Article 9.1.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from AHPND

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.1.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from AHPND, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.1.3. or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of $V_{PAHPND}$ or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of $V_{PAHPND}$ or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.1.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the AHPND status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to AHPND, regardless of the AHPND status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animal products being used for any purpose other than for human consumption.

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.1.2. from a country, zone or compartment not declared free from AHPND, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2017.
CHAPTER 9.2.

INFECTION WITH APHANOMYCES ASTACI
(CRAYFISH PLAGUE)

Article 9.2.1.

For the purposes of the Aquatic Code, infection with Aphanomyces astaci means infection with the pathogenic agent Aphanomyces astaci, of the Family Leptolegniaceae, Phylum Oomycota (water moulds). The disease is commonly known as crayfish plague.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 9.2.2.

Scope

The recommendations in this chapter apply to all species of crayfish in all three crayfish families (Cambaridae, Astacidae and Parastacidae). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 9.2.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with A. astaci status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to A. astaci, regardless of the infection with A. astaci status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.2.2., which are intended for any purpose and comply with Article 5.4.1.:

a) heat sterilised hermetically sealed crayfish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate A. astaci);

b) cooked crayfish products that have been subjected to heat treatment at 100°C for at least one minute (or any time/temperature equivalent that has been demonstrated to inactivate A. astaci);

c) pasteurised crayfish products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent that has been demonstrated to inactivate A. astaci);

d) frozen crayfish products that have been subjected to minus 20°C or lower temperatures for at least 72 hours;

e) crayfish oil;

f) crayfish meal;

g) chemically extracted chitin.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.2.2., other than those referred to in point 1 of Article 9.2.3., Competent Authorities should require the conditions prescribed in Articles 9.2.7. to 9.2.11. relevant to the infection with A. astaci status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.2.2. but which could reasonably be expected to pose a risk of transmission of A. astaci, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.
Article 9.2.4.

Country free from infection with *A. astaci*

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with *A. astaci* if all the areas covered by the shared water bodies are declared countries or zones free from infection with *A. astaci* (see Article 9.2.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with *A. astaci* if:

1) none of the susceptible species referred to in Article 9.2.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.2.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with *A. astaci* for at least the last 25 years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

3) the infection with *A. astaci* status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last five years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last five years without detection of *A. astaci*;

OR

4) it previously made a self-declaration of freedom from infection with *A. astaci* and subsequently lost its free status due to the detection of *A. astaci* but the following conditions have been met:
   a) on detection of *A. astaci*, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of *A. astaci*, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with *A. astaci*; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last five years without detection of *A. astaci*.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.2.5.

Article 9.2.5.

Zone or compartment free from infection with *A. astaci*

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with *A. astaci* if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with *A. astaci* may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.2.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.2.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of infection with *A. astaci* for at least the last 25 years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
b) basic biosecurity conditions have been continuously met for at least the last 10 years;

OR

3) the infection with *A. astaci* status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last five years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last five years without detection of *A. astaci*;

OR

4) it previously made a self-declaration of freedom for a zone from infection with *A. astaci* and subsequently lost its free status due to the detection of *A. astaci* in the zone but the following conditions have been met:
   a) on detection of *A. astaci*, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of *A. astaci*, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with *A. astaci*; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last five years without detection of *A. astaci*.

Article 9.2.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with *A. astaci* following the provisions of points 1 or 2 of Articles 9.2.4. or 9.2.5. (as relevant) may maintain its status as free from infection with *A. astaci* provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with *A. astaci* following the provisions of point 3 of Articles 9.2.4. or 9.2.5. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions are conducive to clinical expression of infection with *A. astaci*, as described in the corresponding chapter of the Aquatic Manual, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with *A. astaci*, targeted surveillance should be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 9.2.7.

Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with *A. astaci*

When importing aquatic animals of a species referred to in Article 9.2.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from infection with *A. astaci*, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country. The international aquatic animal health certificate should state that, on the basis of the procedures described in Articles 9.2.4. or 9.2.5. (as applicable) and 9.2.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from infection with *A. astaci*.

The international aquatic animal health certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.2.3.
Article 9.2.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with A. astaci

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.2.2. from a country, zone or compartment not declared free from infection with A. astaci, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate A. astaci in accordance with Chapters 4.3., 4.7. and 5.5.

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with A. astaci.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for A. astaci in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with A. astaci (as described in Chapter 2.2.2. of the Aquatic Manual) and test for A. astaci in accordance with Chapter 1.4.;
      v) if A. astaci is not detected in the F-1 population, it may be defined as free from infection with A. astaci and may be released from quarantine;
      vi) if A. astaci is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.2.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with A. astaci

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.2.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with A. astaci, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.2.3. or in point 1 of Article 9.2.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of A. astaci or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of A. astaci or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Article 9.2.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with *A. astaci*

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.2.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with *A. astaci*, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.2.3. or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of *A. astaci* or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of *A. astaci* or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.2.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with *A. astaci* status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to *A. astaci* regardless of the infection with *A. astaci* status of the exporting country, zone or compartment, when authorising the importation (or transit) of the following aquatic animal products that have been prepared and packaged for retail trade and comply with Article 5.4.2.:

   – no aquatic animal products listed

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.2.2. from a country, zone or compartment not declared free from infection with *A. astaci*, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 9.3.

INFECTION WITH *HEPATOBACTER PENAEI*

(NECROTISING HEPATOPANCREATEITIS)

Article 9.3.1.

For the purposes of the Aquatic Code, infection with *Hepatobacter penaei* means *infection* with the pathogenic agent *Candidatus Hepatobacter penaei*, an obligate intracellular bacterium of the Order alpha-Proteobacteria. The *disease* is commonly known as necrotising hepatopancreatitis.

Article 9.3.2.

Scope

The recommendations in this chapter apply to the following species that meet the criteria for listing as susceptible in accordance with Chapter 1.5.: whiteleg shrimp (*Penaeus vannamei*).

Article 9.3.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with *H. penaei* status of the exporting country, zone or compartment

1) *Competent Authorities* should not require any conditions related to *H. penaei*, regardless of the infection with *H. penaei* status of the exporting country, zone or compartment, when authorising the importation or transit of the following *aquatic animal products* derived from a species referred to in Article 9.3.2., which are intended for any purpose and comply with Article 5.4.1.:
   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate *H. penaei*);
   b) cooked crustacean products that have been subjected to heat treatment at 100°C for at least three minutes (or any time/temperature equivalent that has been demonstrated to inactivate *H. penaei*);
   c) pasteurised crustacean products that have been subjected to heat treatment at 63°C for at least 30 minutes (or any time/temperature equivalent that has been demonstrated to inactivate *H. penaei*);
   d) crustacean oil;
   e) crustacean meal;
   f) chemically extracted chitin.

2) When authorising the importation or transit of *aquatic animal products* derived from a species referred to in Article 9.3.2., other than those referred to in point 1 of Article 9.3.3., *Competent Authorities* should require the conditions prescribed in Articles 9.3.7. to 9.3.11. relevant to the infection with *H. penaei* status of the exporting country, zone or compartment.

3) When considering the importation or transit of *aquatic animal products* derived from a species not referred to in Article 9.3.2. but which could reasonably be expected to pose a risk of transmission of *H. penaei*, the *Competent Authority* should conduct a *risk analysis* in accordance with the recommendations in Chapter 2.1. The *Competent Authority* of the exporting country should be informed of the outcome of this analysis.

Article 9.3.4.

Country free from infection with *H. penaei*

If a country shares a zone with one or more other countries, it can only make a *self-declaration of freedom* from infection with *H. penaei* if all the areas covered by the shared water bodies are declared countries or zones free from infection with *H. penaei* (see Article 9.3.5.).
Chapter 9.3.- Infection with Hepatobacter penaei

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with *H. penaei* if:

1) none of the susceptible species referred to in Article 9.3.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.3.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with *H. penaei* for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the *Aquatic Manual*); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with *H. penaei* status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of *H. penaei*;

OR

4) it previously made a self-declaration of freedom from infection with *H. penaei* and subsequently lost its free status due to the detection of *H. penaei* but the following conditions have been met:
   a) on detection of *H. penaei*, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of *H. penaei*, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with *H. penaei*; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of *H. penaei*.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.3.5.

Article 9.3.5.

Zone or compartment free from infection with *H. penaei*

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with *H. penaei* if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with *H. penaei* may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.3.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.3.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of infection with *H. penaei* for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the *Aquatic Manual*); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;
3) the infection with \textit{H. penaei} status prior to \textit{targeted surveillance} is unknown but the following conditions have been met:
   
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   
   b) \textit{targeted surveillance}, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of \textit{H. penaei};

OR

4) it previously made a self-declaration of freedom for a zone from infection with \textit{H. penaei} and subsequently lost its free status due to the detection of \textit{H. penaei} in the zone but the following conditions have been met:
   
   a) on detection of \textit{H. penaei}, the affected area was declared an \textit{infected zone} and a \textit{protection zone} was established; and
   
   b) infected populations within the \textit{infected zone} have been killed and disposed of by means that minimise the likelihood of further transmission of \textit{H. penaei}, and the appropriate \textit{disinfection} procedures (as described in Chapter 4.3.) have been completed; and
   
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with \textit{H. penaei}; and
   
   d) \textit{targeted surveillance}, as described in Chapter 1.4., has been in place for at least the last two years without detection of \textit{H. penaei}.

\textbf{Article 9.3.6.}

\textbf{Maintenance of free status}

A country, zone or compartment that is declared free from infection with \textit{H. penaei} following the provisions of points 1 or 2 of Articles 9.3.4. or 9.3.5. (as relevant) may maintain its status as free from infection with \textit{H. penaei} provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with \textit{H. penaei} following the provisions of point 3 of Articles 9.3.4. or 9.3.5. (as relevant) may discontinue \textit{targeted surveillance} and maintain its free status provided that conditions are conducive to clinical expression of infection with \textit{H. penaei}, as described in the corresponding chapter of the \textit{Aquatic Manual}, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with \textit{H. penaei}, \textit{targeted surveillance} should be continued at a level determined by the \textit{Aquatic Animal Health Service} on the basis of the likelihood of infection.

\textbf{Article 9.3.7.}

\textbf{Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with \textit{H. penaei}}

When importing aquatic animals of a species referred to in Article 9.3.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from infection with \textit{H. penaei}, the \textit{Competent Authority} of the \textit{importing country} should require that the consignment be accompanied by an \textit{international aquatic animal health certificate} issued by the \textit{Competent Authority} of the \textit{exporting country} or a \textit{certifying official} approved by the \textit{importing country}. The \textit{international aquatic animal health certificate} should state that, on the basis of the procedures described in Articles 9.3.4. or 9.3.5. (as applicable) and 9.3.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from infection with \textit{H. penaei}.

The \textit{international aquatic animal health certificate} should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.3.3.
Chapter 9.3.- Infection with Hepatobacter penaei

Article 9.3.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with *H. penaei*

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.3.2. from a country, zone or compartment not declared free from infection with *H. penaei*, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate *H. penaei* in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with *H. penaei*.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for *H. penaei* in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with *H. penaei* (as described in Chapter 2.2.3. of the Aquatic Manual) and test for *H. penaei* in accordance with Chapter 1.4.;
      v) if *H. penaei* is not detected in the F-1 population, it may be defined as free from infection with *H. penaei* and may be released from quarantine;
      vi) if *H. penaei* is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.3.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with *H. penaei*

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.3.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with *H. penaei*, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.3.3. or in point 1 of Article 9.3.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of *H. penaei* or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of *H. penaei* or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Chapter 9.3.- Infection with Hepatobacter penaei

Article 9.3.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with H. penaei

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.3.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with H. penaei, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.3.3. or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of H. penaei or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of H. penaei or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.3.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with H. penaei status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to H. penaei, regardless of the infection with H. penaei status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animal products being used for any purpose other than for human consumption.

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.3.2. from a country, zone or compartment not declared free from infection with H. penaei, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2010; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 9.4.

INFECTION WITH INFECTIOUS HYPODERMAL AND HAEMATOPOIETIC NECROSIS VIRUS

Article 9.4.1.

For the purposes of the Aquatic Code, infection with infectious hypodermal and haematopoietic necrosis virus means infection with the pathogenic agent infectious hypodermal and haematopoietic necrosis virus (IHHNV), of the Genus Brevidensovirus and Family Paroviridae.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 9.4.2.

Scope

The recommendations in this chapter apply to the following species that meet the criteria for listing as susceptible in accordance with Chapter 1.5.: giant river prawn (Macrobrachium rosenbergii) (under study), yellowleg shrimp (Penaeus californiensis), giant tiger prawn (Penaeus monodon), northern white shrimp (Penaeus setiferus), blue shrimp (Penaeus stylirostris) and whiteleg shrimp (Penaeus vannamei).

Article 9.4.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with IHHNV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to IHHNV regardless of the infection with IHHNV status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.4.2., which are intended for any purpose and comply with Article 5.4.1.:

   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate IHHNV);

   b) cooked crustacean products that have been subjected to heat treatment at 90°C for at least 20 minutes (or any time/temperature equivalent that has been demonstrated to inactivate IHHNV);

   c) crustacean oil;

   d) crustacean meal.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.4.2., other than those referred to in point 1 of Article 9.4.3., Competent Authorities should require the conditions prescribed in Articles 9.4.7. to 9.4.11. relevant to the infection with IHHNV status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.4.2. but which could reasonably be expected to pose a risk of transmission of IHHNV, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.

Article 9.4.4.

Country free from infection with IHHNV

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with IHHNV if all the areas covered by the shared water bodies are declared countries or zones free from infection with IHHNV (see Article 9.4.5.).
As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with IHHNV if:

1) none of the susceptible species referred to in Article 9.4.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.4.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with IHHNV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with IHHNV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IHHNV;

OR

4) it previously made a self-declaration of freedom from infection with IHHNV and subsequently lost its free status due to the detection of IHHNV but the following conditions have been met:
   a) on detection of IHHNV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of IHHNV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with IHHNV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IHHNV.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.4.5.

Article 9.4.5.

Zone or compartment free from infection with IHHNV

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with IHHNV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with IHHNV may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.4.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.4.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of infection with IHHNV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;
Chapter 9.4.- Infection with infectious hypodermal and haematopoietic necrosis virus

OR

3) the infection with IHHNV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of IHHNV;

OR

4) it previously made a self-declaration of freedom for a zone from infection with IHHNV and subsequently lost its free status due to the detection of IHHNV in the zone but the following conditions have been met:
   a) on detection of IHHNV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of IHHNV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with IHHNV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IHHNV.

Article 9.4.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with IHHNV following the provisions of points 1 or 2 of Articles 9.4.4. or 9.4.5. (as relevant) may maintain its status as free from infection with IHHNV provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with IHHNV following the provisions of point 3 of Articles 9.4.4. or 9.4.5. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions are conducive to clinical expression of infection with IHHNV, as described in the corresponding chapter of the Aquatic Manual, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with IHHNV, targeted surveillance should be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 9.4.7.

Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with IHHNV

When importing aquatic animals of a species referred to in Article 9.4.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from infection with IHHNV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country. The international aquatic animal health certificate should state that, on the basis of the procedures described in Articles 9.4.4. or 9.4.5. (as applicable) and 9.4.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from infection with IHHNV.

The international aquatic animal health certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.4.3.
Article 9.4.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with IHHNV

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.4.2., from a country, zone or compartment not declared free from infection with IHHNV, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate IHHNV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with IHHNV.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for IHHNV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with IHHNV (as described in Chapter 2.2.4. of the Aquatic Manual) and test for IHHNV in accordance with Chapter 1.4.;
      v) if IHHNV is not detected in the F-1 population, it may be defined as free from infection with IHHNV and may be released from quarantine;
      vi) if IHHNV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.4.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with IHHNV

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.4.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with IHHNV, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.4.3. or in point 1 of Article 9.4.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of IHHNV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of IHHNV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Article 9.4.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with IHHNV

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, *aquatic animals* of a species referred to in Article 9.4.2., or *aquatic animal products* derived thereof, from a country, zone or compartment not declared free from infection with IHHNV, the *Competent Authority* of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.4.3. or other products authorised by the *Competent Authority*; and

2) all containers and water used in transport are treated to ensure inactivation of IHHNV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of IHHNV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.4.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with IHHNV status of the exporting country, zone or compartment

1) *Competent Authorities* should not require any conditions related to IHHNV, regardless of the infection with IHHNV status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2. Certain assumptions have been made in assessing the safety of the *aquatic animal products* mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these *aquatic animal products* Member Countries may wish to consider introducing internal measures to address the *risks* associated with the *aquatic animal products* being used for any purpose other than for human consumption.

2) When importing *aquatic animal products*, other than those referred to in point 1 above, derived from a species referred to in Article 9.4.2. from a country, zone or compartment not declared free from infection with IHHNV, the *Competent Authority* of the importing country should assess the *risk* and apply appropriate *risk* mitigation measures.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 9.5.

INFECTION WITH INFECTIOUS MYONECROSIS VIRUS

Article 9.5.1.

For the purposes of the Aquatic Code, infection with infectious myonecrosis virus means infection with the pathogenic agent infectious myonecrosis virus (IMNV) of the Family Totiviridae (tentative classification).

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 9.5.2.

Scope

The recommendations in this chapter apply to the following species that meet the criteria for listing as susceptible in accordance with Chapter 1.5.: brown tiger prawn (Penaeus esculentus), banana prawn (Penaeus merguiensis) and whiteleg shrimp (Penaeus vannamei).

Article 9.5.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with IMNV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to IMNV, regardless of the infection with IMNV status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.5.2., which are intended for any purpose and comply with Article 5.4.1.:

   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate IMNV);
   b) cooked crustacean products that have been subjected to heat treatment at 60°C for at least three minutes (or any time/temperature equivalent that has been demonstrated to inactivate IMNV);
   c) crustacean oil;
   d) crustacean meal;
   e) chemically extracted chitin.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.5.2., other than those referred to in point 1 of Article 9.5.3., Competent Authorities should require the conditions prescribed in Articles 9.5.7. to 9.5.11. relevant to the infection with IMNV status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.5.2. but which could reasonably be expected to pose a risk of transmission of IMNV, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.

Article 9.5.4.

Country free from infection with IMNV

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with IMNV if all the areas covered by the shared water bodies are declared countries or zones free from infection with IMNV (see Article 9.5.5.).
Chapter 9.5.- Infection with infectious myonecrosis virus

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with IMNV if:

1) none of the susceptible species referred to in Article 9.5.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.5.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with IMNV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with IMNV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IMNV;

OR

4) it previously made a self-declaration of freedom from infection with IMNV and subsequently lost its free status due to the detection of IMNV but the following conditions have been met:
   a) on detection of IMNV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of IMNV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with IMNV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IMNV.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.5.5.

Article 9.5.5.

Zone or compartment free from infection with IMNV

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with IMNV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with IMNV may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.5.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.5.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of infection with IMNV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with IMNV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of IMNV;
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OR

4) it previously made a self-declaration of freedom for a zone from infection with IMNV and subsequently lost its free status due to the detection of IMNV in the zone but the following conditions have been met:

a) on detection of IMNV, the affected area was declared an infected zone and a protection zone was established; and

b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of IMNV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and

c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with IMNV; and

d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IMNV.

Article 9.5.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with IMNV following the provisions of points 1 or 2 of Articles 9.5.4. or 9.5.5. (as relevant) may maintain its status as free from infection with IMNV provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with IMNV following the provisions of point 3 of Articles 9.5.4. or 9.5.5. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions are conducive to clinical expression of infection with IMNV as described in the corresponding chapter of the Aquatic Manual, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with IMNV, targeted surveillance should be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 9.5.7.

Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with IMNV

When importing aquatic animals of a species referred to in Article 9.5.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from infection with IMNV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country. The international aquatic animal health certificate should state that, on the basis of the procedures described in Articles 9.5.4. or 9.5.5. (as applicable) and 9.5.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from infection with IMNV.

The international aquatic animal health certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.5.3.
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Article 9.5.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with IMNV

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.5.2. from a country, zone or compartment not declared free from infection with IMNV, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate IMNV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with IMNV.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for IMNV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with IMNV (as described in Chapter 2.2.5. of the Aquatic Manual) and test for IMNV in accordance with Chapter 1.4.;
      v) if IMNV is not detected in the F-1 population, it may be defined as free from infection with IMNV and may be released from quarantine;
      vi) if IMNV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.5.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with IMNV

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.5.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with IMNV, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.5.3. or in point 1 of Article 9.5.11., or other products authorised by the Competent Authority; and
2) all containers and water used in transport are treated to ensure inactivation of IMNV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and
3) all processing effluent and waste materials are treated to ensure inactivation of IHHNV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
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Article 9.5.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with IMNV

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.5.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with IMNV, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.5.3. or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of IMNV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of IMNV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.5.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with IMNV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to IMNV, regardless of the infection with IMNV status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2. Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animal products being used for any purpose other than for human consumption.

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.5.2. from a country, zone or compartment not declared free from infection with IMNV, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2008; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 9.6.

INFECTION WITH MACROBRACHIUM ROSENBERGII NODAVIRUS (WHITE TAIL DISEASE)

Article 9.6.1.

For the purposes of the Aquatic Code, infection with Macrobrachium rosenbergii nodavirus means infection with the pathogenic agent Macrobrachium rosenbergii nodavirus (MrNV), of the Family Nodaviridae. The disease is commonly known as white tail disease.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 9.6.2.

Scope

The recommendations in this chapter apply to the following species that meet the criteria for listing as susceptible in accordance with Chapter 1.5.: giant river prawn (Macrobrachium rosenbergii).

Article 9.6.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with MrNV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to MrNV, regardless of the infection with MrNV status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.6.2., which are intended for any purpose and comply with Article 5.4.1.:
   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate MrNV);
   b) cooked crustacean products that have been subjected to heat treatment at 60°C for at least 60 minutes (or any time/temperature equivalent that has been demonstrated to inactivate MrNV);
   c) pasteurised crustacean products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent that has been shown to inactivate MrNV);
   d) crustacean oil;
   e) crustacean meal;
   f) chemically extracted chitin.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.6.2., other than those referred to in point 1 of Article 9.6.3., Competent Authorities should require the conditions prescribed in Articles 9.6.7. to 9.6.11. relevant to the infection with MrNV status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.6.2. but which could reasonably be expected to pose a risk of transmission of MrNV, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.
Chapter 9.6. Infection with Macrobrachium rosenbergii nodavirus

Article 9.6.4.

Country free from infection with MrNV

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with MrNV if all the areas covered by the shared water bodies are declared countries or zones free from infection with MrNV (see Article 9.6.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with MrNV if:

1) none of the susceptible species referred to in Article 9.6.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.6.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with MrNV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with MrNV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of MrNV;

OR

4) it previously made a self-declaration of freedom from infection with MrNV and subsequently lost its free status due to the detection of MrNV but the following conditions have been met:
   a) on detection of MrNV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of MrNV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with MrNV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of MrNV.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.6.5.

Article 9.6.5.

Zone or compartment free from infection with MrNV

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with MrNV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with MrNV may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.6.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.6.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of infection with MrNV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;
Chapter 9.6. - Infection with Macrobrachium rosenbergii nodavirus

OR

3) the infection with MrNV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of MrNV;

OR

4) it previously made a self-declaration of freedom for a zone from infection with MrNV and subsequently lost its free status due to the detection of MrNV in the zone but the following conditions have been met:
   a) on detection of MrNV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of MrNV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with MrNV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of MrNV.

Article 9.6.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with MrNV following the provisions of points 1 or 2 of Articles 9.6.4. or 9.6.5. (as relevant) may maintain its status as free from infection with MrNV provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with MrNV following the provisions of point 3 of Articles 9.6.4. or 9.6.5. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions are conducive to clinical expression of infection with MrNV, as described in the corresponding chapter of the Aquatic Manual, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with MrNV, targeted surveillance should be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 9.6.7.

Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with MrNV

When importing aquatic animals of a species referred to in Article 9.6.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from infection with MrNV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country. The international aquatic animal health certificate should state that, on the basis of the procedures described in Articles 9.6.4. or 9.6.5. (as applicable) and 9.6.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from infection with MrNV.

The international aquatic animal health certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.6.3.
Chapter 9.6.- Infection with Macrobrachium rosenbergii nodavirus

Article 9.6.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with MrNV

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.6.2. from a country, zone or compartment not declared free from infection with MrNV, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate MrNV in accordance with Chapters 4.3., 4.7. and 5.5.

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with MrNV.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for MrNV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with MrNV (as described in Chapter 2.2.6. of the Aquatic Manual) and test for MrNV in accordance with Chapter 1.4.;
      v) if MrNV is not detected in the F-1 population, it may be defined as free from infection with MrNV and may be released from quarantine;
      vi) if MrNV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.6.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with MrNV

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.6.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with MrNV, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.6.3. or in point 1 of Article 9.6.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of MrNV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of MrNV or disposed of in a biosecure manner in accordance with Chapters 4.3 and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Chapter 9.6. - Infection with Macrobrachium rosenbergii nodavirus

Article 9.6.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with MrNV

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.6.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with MrNV, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.6.3. or other products authorised by the Competent Authority; and
2) all containers and water used in transport are treated to ensure inactivation of MrNV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and
3) all processing effluent and waste materials are treated to ensure inactivation of MrNV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.6.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with MrNV status of the exporting country, zone or compartment.

1) Competent Authorities should not require any conditions related to MrNV, regardless of the infection with MrNV status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2. Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animal products being used for any purpose other than for human consumption.

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.6.2. from a country, zone or compartment not declared free from infection with MrNV, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2008; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 9.7.

INFECTION WITH TAURA SYNDROME VIRUS

Article 9.7.1.

For the purposes of the Aquatic Code, infection with Taura syndrome virus means infection with the pathogenic agent Taura syndrome virus (TSV), of the Genus Aparavirus, Family Dicistroviridae and Order Picornavirales.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 9.7.2.

Scope

The recommendations in this chapter apply to the following species that meet the criteria for listing as susceptible in accordance with Chapter 1.5.: greasyback shrimp (Metapenaeus ensis), northern brown shrimp (Penaeus aztecus), giant tiger prawn (Penaeus monodon), northern white shrimp (Penaeus setiferus), blue shrimp (Penaeus stylirostris) and whiteleg shrimp (Penaeus vannamei).

Article 9.7.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with TSV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to TSV, regardless of the infection with TSV status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.7.2., which are intended for any purpose and comply with Article 5.4.1.:

   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate TSV);
   b) cooked crustacean products that have been subjected to heat treatment at 70°C for at least 30 minutes (or any time/temperature equivalent that has been demonstrated to inactivate TSV);
   c) pasteurised crustacean products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent that has been demonstrated to inactivate TSV);
   d) crustacean oil;
   e) crustacean meal;
   f) chemically extracted chitin.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.7.2., other than those referred to in point 1 of Article 9.7.3., Competent Authorities should require the conditions prescribed in Articles 9.7.7. to 9.7.11. relevant to the infection with TSV status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.7.2. but which could reasonably be expected to pose a risk of transmission of TSV, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.
Chapter 9.7.- Infection with Taura syndrome virus

Article 9.7.4.

Country free from infection with TSV

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with TSV if all the areas covered by the shared water bodies are declared countries or zones free from infection with TSV (see Article 9.7.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with TSV if:

1) none of the susceptible species referred to in Article 9.7.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.7.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with TSV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with TSV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of TSV;

OR

4) it previously made a self-declaration of freedom from infection with TSV and subsequently lost its free status due to the detection of TSV but the following conditions have been met:
   a) on detection of TSV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of TSV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with TSV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of TSV.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.7.5.

Article 9.7.5.

Zone or compartment free from infection with TSV

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with TSV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with TSV may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.7.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.7.2. are present in the zone or compartment and the following conditions have been made:
   a) there has not been any occurrence of infection with TSV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;
Chapter 9.7. Infection with Taura syndrome virus

OR

3) the infection with TSV status prior to targeted surveillance is unknown but the following conditions have been made:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of TSV;

OR

4) it previously made a self-declaration of freedom for a zone from infection with TSV and subsequently lost its free status due to the detection of TSV in the zone but the following conditions have been met:
   a) on detection of TSV the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of TSV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with TSV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of TSV.

Article 9.7.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with TSV following the provisions of points 1 or 2 of Articles 9.7.4. or 9.7.5. (as relevant) may maintain its status as free from infection with TSV provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with TSV following the provisions of point 3 of Articles 9.7.4. or 9.7.5. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions are conducive to clinical expression of infection with TSV, as described in the corresponding chapter of the Aquatic Manual, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with TSV, targeted surveillance should be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 9.7.7.

Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with TSV

When importing aquatic animals of a species referred to in Article 9.7.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from infection with TSV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country. The international aquatic animal health certificate should state that, on the basis of the procedures described in Articles 9.6.4. or 9.6.5. (as applicable) and 9.6.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from infection with TSV.

The international aquatic animal health certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.7.3.
Chapter 9.7.- Infection with Taura syndrome virus

Article 9.7.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with TSV

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.7.2. from a country, zone or compartment not declared free from infection with TSV, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate TSV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with TSV.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for TSV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with TSV (as described in Chapter 2.2.7. of the Aquatic Manual) and test for TSV in accordance with Chapter 1.4.;
      v) if TSV is not detected in the F-1 population, it may be defined as free from infection with TSV and may be released from quarantine;
      vi) if TSV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.7.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with TSV

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.7.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with TSV, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.7.3. or in point 1 of Article 9.7.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of TSV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of TSV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Chapter 9.7. - Infection with Taura syndrome virus

Article 9.7.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with TSV

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.7.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with TSV, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.7.3. or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of TSV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of TSV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.7.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with TSV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to TSV, regardless of the infection with TSV status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp or decapod crustacea (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animal products being used for any purpose other than for human consumption.

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.7.2. from a country, zone or compartment not declared free from infection with TSV, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 9.8.

INFECTION WITH WHITE SPOT SYNDROME VIRUS

Article 9.8.1.

For the purposes of the Aquatic Code, infection with white spot syndrome virus means infection with the pathogenic agent white spot syndrome virus (WSSV), of the Genus Whispovirus, Family Nimaviridae.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 9.8.2.

Scope

The recommendations in this chapter apply to all decapod (order Decapoda) crustaceans from marine, brackish and freshwater sources. These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 9.8.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with WSSV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to WSSV, regardless of the infection with WSSV status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.8.2., which are intended for any purpose and comply with Article 5.4.1.:
   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate WSSV);
   b) cooked crustacean products that have been subjected to heat treatment at 60°C for at least one minute (or any time/temperature equivalent that has been demonstrated to inactivate WSSV);
   c) pasteurised crustacean products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent that has been demonstrated to inactivate WSSV);
   d) crustacean oil;
   e) crustacean meal;
   f) chemically extracted chitin.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.8.2., other than those referred to in point 1 of Article 9.8.3., Competent Authorities should require the conditions prescribed in Articles 9.8.7. to 9.8.11. relevant to the infection with WSSV status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.8.2. but which could reasonably be expected to pose a risk of transmission of WSSV, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.

Article 9.8.4.

Country free from infection with WSSV

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with WSSV if all the areas covered by the shared water bodies are declared countries or zones free from infection with WSSV (see Article 9.8.5.).
As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with WSSV if:

1) none of the susceptible species referred to in Article 9.8.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.8.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with WSSV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with WSSV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of WSSV;

OR

4) it previously made a self-declaration of freedom from infection with WSSV and subsequently lost its free status due to the detection of WSSV but the following conditions have been met:
   a) on detection of WSSV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of WSSV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with WSSV; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of WSSV.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.8.5.

Article 9.8.5.

Zone or compartment free from infection with WSSV

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with WSSV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with WSSV may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.8.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.8.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of infection with WSSV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with WSSV status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of WSSV;
OR

4) it previously made a *self-declaration of freedom* for a *zone* from infection with WSSV and subsequently lost its free status due to the detection of WSSV in the *zone* but the following conditions have been met:

   a) on detection of WSSV, the affected area was declared an *infected zone* and a *protection zone* was established; and

   b) infected populations within the *infected zone* have been killed and disposed of by means that minimise the likelihood of further transmission of WSSV, and the appropriate *disinfection* procedures (as described in Chapter 4.3.) have been completed; and

   c) previously existing *basic biosecurity conditions* have been reviewed and modified as necessary and have continuously been in place since eradication of infection with WSSV; and

   d) *targeted surveillance*, as described in Chapter 1.4., has been in place for at least the last two years without detection of WSSV.

**Article 9.8.6.**

**Maintenance of free status**

A country, *zone* or *compartment* that is declared free from infection with WSSV following the provisions of points 1 or 2 of Articles 9.8.4. or 9.8.5. (as relevant) may maintain its status as free from infection with WSSV provided that *basic biosecurity conditions* are continuously maintained.

A country, *zone* or *compartment* that is declared free from infection with WSSV following the provisions of point 3 of Articles 9.8.4. or 9.8.5. (as relevant) may discontinue *targeted surveillance* and maintain its free status provided that conditions are conducive to clinical expression of infection with WSSV, as described in the corresponding chapter of the *Aquatic Manual*, and *basic biosecurity conditions* are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with WSSV, *targeted surveillance* should be continued at a level determined by the *Aquatic Animal Health Service* on the basis of the likelihood of infection.

**Article 9.8.7.**

**Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with WSSV**

When importing *aquatic animals* of a species referred to in Article 9.8.2., or *aquatic animal products* derived thereof, from a country, *zone* or *compartment* declared free from infection with WSSV, the *Competent Authority* of the *importing country* should require that the consignment be accompanied by an *international aquatic animal health certificate* issued by the *Competent Authority* of the *exporting country* or a *certifying official* approved by the *importing country*. The *international aquatic animal health certificate* should state that, on the basis of the procedures described in Articles 9.8.4. or 9.8.5. (as applicable) and 9.8.6., the place of production of the *aquatic animals* or *aquatic animal products* is a country, *zone* or *compartment* declared free from infection with WSSV.

The *international aquatic animal health certificate* should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to *aquatic animal products* listed in point 1 of Article 9.8.3.
Article 9.8.8.

**Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with WSSV**

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.8.2. from a country, zone or compartment not declared free from infection with WSSV, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate WSSV in accordance with Chapters 4.3., 4.7. and 5.5.

 OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with WSSV.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for WSSV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with WSSV (as described in Chapter 2.2.8. of the Aquatic Manual) and test for WSSV in accordance with Chapter 1.4.;
      v) if WSSV is not detected in the F-1 population, it may be defined as free from infection with WSSV and may be released from quarantine;
      vi) if WSSV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.8.9.

**Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with WSSV**

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.8.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with WSSV, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held, in quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.8.3. or in point 1 of Article 9.8.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of WSSV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of WSSV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Chapter 9.8.- Infection with white spot syndrome virus

Article 9.8.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with WSSV

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.8.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with WSSV, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.8.3. or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of WSSV or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of WSSV or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.8.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with WSSV status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to WSSV, regardless of the infection with WSSV status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp or decapod crustacea (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animal products being used for any purpose other than for human consumption.

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.8.2. from a country, zone or compartment not declared free from infection with WSSV, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 1997; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 9.9.

INFECTION WITH YELLOW HEAD VIRUS GENOTYPE 1

Article 9.9.1.

For the purposes of the Aquatic Code, infection with yellow head virus genotype 1 means infection with the pathogenic agent yellow head virus genotype 1 (YHV1), of the Genus Okavirus, Family Roniviridae, Order Nidovirales.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 9.9.2.

Scope

The recommendations in this chapter apply to the following species that meet the criteria for listing as susceptible in accordance with Chapter 1.5.: Jinga shrimp (Metapenaeus affinis), giant tiger prawn (Penaeus monodon), dagger blade grass shrimp (Palaemonetes pugio), blue shrimp (Penaeus stylirostris) and whiteleg shrimp (Penaeus vannamei).

Article 9.9.3.

Importation or transit of aquatic animal products for any purpose regardless of the infection with YHV1 status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to YHV1, regardless of the infection with YHV1 status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products derived from a species referred to in Article 9.9.2., which are intended for any purpose and comply with Article 5.4.1.:
   a) heat sterilised hermetically sealed crustacean products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent that has been demonstrated to inactivate YHV1);
   b) cooked crustacean products that have been subjected to heat treatment at 60°C for at least 15 minutes (or any time/temperature equivalent that has been demonstrated to inactivate YHV1);
   c) pasteurised crustacean products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent that has been demonstrated to inactivate YHV1);
   d) crustacean oil;
   e) crustacean meal;
   f) chemically extracted chitin.

2) When authorising the importation or transit of aquatic animal products derived from a species referred to in Article 9.9.2., other than those referred to in point 1 of Article 9.9.3., Competent Authorities should require the conditions prescribed in Articles 9.9.7. to 9.9.11. relevant to the infection with YHV1 status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animal products derived from a species not referred to in Article 9.9.2. but which could reasonably be expected to pose a risk of transmission of YHV1, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this analysis.
Article 9.9.4.

Country free from infection with yellow head virus genotype 1

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with YHV1 if all the areas covered by the shared water bodies are declared countries or zones free from infection with YHV1 (see Article 9.9.5).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with YHV1 if:

1) none of the susceptible species referred to in Article 9.9.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.9.2. are present and the following conditions have been met:
   a) there has been no occurrence of infection with YHV1 for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the infection with YHV1 status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of YHV1;

OR

4) it previously made a self-declaration of freedom from infection with YHV1 and subsequently lost its free status due to the detection of YHV1 but the following conditions have been met:
   a) on detection of YHV1, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of YHV1, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with YHV1; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of YHV1.

In the meantime, part or all of the unaffected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 9.9.5.

Article 9.9.5.

Zone or compartment free from infection with yellow head virus genotype 1

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with YHV1 if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with YHV1 may be declared free by the Competent Authority of the country concerned if:

1) none of the susceptible species referred to in Article 9.9.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 9.9.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any occurrence of infection with YHV1 for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;
OR

3) the infection with YHV1 status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of YHV1;

OR

4) it previously made a self-declaration of freedom for a zone from infection with YHV1 and subsequently lost its free status due to the detection of YHV1 in the zone but the following conditions have been met:
   a) on detection of YHV1, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations within the infected zone have been killed and disposed of by means that minimise the likelihood of further transmission of YHV1, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of infection with YHV1; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of YHV1.

Article 9.9.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with YHV1 following the provisions of points 1 or 2 of Articles 9.9.4. or 9.9.5. (as relevant) may maintain its status as free from infection with YHV1 provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with YHV1 following the provisions of point 3 of Articles 9.9.4. or 9.9.5. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions are conducive to clinical expression of infection with YHV1, as described in the corresponding chapter of the Aquatic Manual, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with YHV1, targeted surveillance should be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 9.9.7.

Importation of aquatic animals or aquatic animal products from a country, zone or compartment declared free from infection with yellow head virus genotype 1

When importing aquatic animals of a species referred to in Article 9.9.2., or aquatic animal products derived thereof, from a country, zone or compartment declared free from infection with YHV1, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country. The international aquatic animal health certificate should state that, on the basis of the procedures described in Articles 9.9.4. or 9.9.5. (as applicable) and 9.9.6., the place of production of the aquatic animals or aquatic animal products is a country, zone or compartment declared free from infection with YHV1.

The international aquatic animal health certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to aquatic animal products listed in point 1 of Article 9.9.3.
Chapter 9.9. - Infection with yellow head virus genotype 1

Article 9.9.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with yellow head virus genotype 1

When importing, for aquaculture, aquatic animals of a species referred to in Article 9.9.2. from a country, zone or compartment not declared free from infection with YHV1, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of transport water, equipment, effluent and waste materials to inactivate YHV1 in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following.
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with YHV1.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for YHV1 in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with YHV1 (as described in Chapter 2.2.9. of the Aquatic Manual) and test for YHV1 in accordance with Chapter 1.4.;
      v) if YHV1 is not detected in the F-1 population, it may be defined as free from infection with YHV1 and may be released from quarantine;
      vi) if YHV1 is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 9.9.9.

Importation of aquatic animals or aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with yellow head virus genotype 1

When importing, for processing for human consumption, aquatic animals of a species referred to in Article 9.9.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with YHV1, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.9.3. or in point 1 of Article 9.9.11., or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of YHV1 or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of YHV1 or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

For these aquatic animals or aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animals or aquatic animal products being used for any purpose other than for human consumption.
Article 9.9.10.

Importation of aquatic animals or aquatic animal products intended for uses other than human consumption including animal feed, or for agricultural, industrial, research or pharmaceutical use, from a country, zone or compartment not declared free from infection with yellow head virus genotype 1

When importing, for use in animal feed or for agricultural, industrial, research or pharmaceutical use, aquatic animals of a species referred to in Article 9.9.2., or aquatic animal products derived thereof, from a country, zone or compartment not declared free from infection with YHV1, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processed into one of the products referred to in point 1 of Article 9.9.3. or other products authorised by the Competent Authority; and

2) all containers and water used in transport are treated to ensure inactivation of YHV1 or disposed of in a biosecure manner in accordance with Chapters 4.3., 4.7. and 5.5.; and

3) all processing effluent and waste materials are treated to ensure inactivation of YHV1 or disposed of in a biosecure manner in accordance with Chapters 4.3. and 4.7.

Article 9.9.11.

Importation (or transit) of aquatic animal products for retail trade for human consumption regardless of the infection with yellow head virus genotype 1 status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to YHV1, regardless of the infection with YHV1 status of the exporting country, zone or compartment, when authorising the importation (or transit) of frozen peeled shrimp or decapod crustacea (shell off, head off) that have been prepared and packaged for retail trade and comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these aquatic animal products Member Countries may wish to consider introducing internal measures to address the risks associated with the aquatic animal products being used for any purpose other than for human consumption.

2) When importing aquatic animal products, other than those referred to in point 1 above, derived from a species referred to in Article 9.9.2. from a country, zone or compartment not declared free from infection with YHV1, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2017.
SECTION 10.

DISEASES OF FISH

CHAPTER 10.1.

EPIZOOTIC HAEMATOPOIETIC NECROSIS

Article 10.1.1.

For the purposes of the Aquatic Code, epizootic haematopoietic necrosis (EHN) means infection with EHN virus (EHNV) of the genus Ranavirus of the family Iridoviridae.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 10.1.2.

Scope

The recommendations in this chapter apply to: redfin perch (Perca fluviatilis) and rainbow trout (Oncorhynchus mykiss). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.1.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the epizootic haematopoietic necrosis status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to EHN, regardless of the EHN status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.1.2. which are intended for any purpose and which comply with Article 5.4.1.:

a) heat sterilised hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);

b) pasteurised fish products that have been subjected to heat treatment at 90°C for ten minutes (or any time/temperature equivalent which has been demonstrated to inactivate EHNV);

c) mechanically dried eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate EHNV);

d) fish oil;

e) fish meal;

f) fish skin leather.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of species referred to in Article 10.1.2., other than those referred to in point 1 of Article 10.1.3., Competent Authorities should require the conditions prescribed in Articles 10.1.7. to 10.1.11. relevant to the EHN status of the exporting country, zone or compartment.
3) When considering the importation or transit of aquatic animals and aquatic animal products of species not covered in Article 10.1.2. but which could reasonably be expected to pose a risk of spread of EHN, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 10.1.4.

Country free from epizootic haematopoietic necrosis

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from EHN if all the areas covered by the shared water bodies are declared countries or zones free from EHN (see Article 10.1.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from EHN if:

1) none of the susceptible species referred to in Article 10.1.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.1.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the past ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of EHN;

OR

4) it previously made a self-declaration of freedom from EHN and subsequently lost its disease free status due to the detection of EHN but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of EHN.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.1.5.

Article 10.1.5.

Zone or compartment free from epizootic haematopoietic necrosis

If a zone or compartment extends over more than one country, it can only be declared an EHN free zone or compartment if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from EHN may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 10.1.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;
OR

2) any of the susceptible species referred to in Article 10.1.2. are present in the zone or compartment and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of EHN;

OR

4) it previously made a self-declaration of freedom for a zone from EHN and subsequently lost its disease free status due to the detection of EHN in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of EHN.

Article 10.1.6.

Maintenance of free status

A country, zone or compartment that is declared free from EHN following the provisions of points 1 or 2 of Articles 10.1.4. or 10.1.5. (as relevant) may maintain its status as free from EHN provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from EHN following the provisions of point 3 of Articles 10.1.4. or 10.1.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from EHN provided that conditions that are conducive to clinical expression of EHN, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of EHN, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.1.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from epizootic haematopoietic necrosis

When importing aquatic animals and aquatic animal products of species referred to in Article 10.1.2. from a country, zone or compartment declared free from EHN, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.1.4. or 10.1.5. (as applicable) and 10.1.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from EHN.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.1.3.
Chapter 10.1.- Epizootic haematopoietic necrosis

Article 10.1.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from epizootic haematopoietic necrosis

When importing for aquaculture, aquatic animals of a species referred to in Article 10.1.2. from a country, zone or compartment not declared free from EHN, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive EHNV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for EHN.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for EHNV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of EHN (as described in Chapter 2.3.1. of the Aquatic Manual) and test for EHNV in accordance with Chapter 1.4.;
      v) if EHNV is not detected in the F-1 population, it may be defined as free from EHN and may be released from quarantine;
      vi) if EHNV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.1.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from epizootic haematopoietic necrosis

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.1.2. from a country, zone or compartment not declared free from EHN, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.1.3., or products described in point 1 of Article 10.1.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of EHNV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 10.1. - Epizootic haematopoietic necrosis

Article 10.1.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from epizootic haematopoietic necrosis

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.1.2. from a country, zone or compartment not declared free from EHN, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held, in quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of EHNV.

This article does not apply to commodities referred to in point 1 of Article 10.1.3.

Article 10.1.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from epizootic haematopoietic necrosis

1) Competent Authorities should not require any conditions related to EHN, regardless of the EHN status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (chilled or frozen) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.1.2. from a country, zone or compartment not declared free from EHN, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.2.

INFECTION WITH APHANOMYCES INVADANS
(EPIZOOTIC ULCERATIVE SYNDROME)

Article 10.2.1.

For the purposes of the Aquatic Code, infection with Aphanomyces invadans means all infections caused by the Oomycete fungus A. invadans (syn. A. piscicida). The disease was previously referred to as epizootic ulcerative syndrome.

Standards for diagnostic tests are described in the Aquatic Manual.

Article 10.2.2.

Scope

The recommendations in this chapter apply to: yellowfin seabream (Acanthopagrus australis), climbing perch (Anabas testudineus), eels (Anguillidae), bagrid catfishes (Bagridae), silver perch (Bidyanus bidyanus), Atlantic menhaden (Brevortia tyrannus), jacks (Caranx spp.), catla (Catla catla), striped snakehead (Channa striatus), mrigal (Cirrhinus mrigala), torpedo-shaped catfishes (Clarius spp.), halfbeaks flying fishes (Exocoetidae), tank goby (Glossogobius giuris), marble goby (Oxyeleotris marmoratus), gobies (Gobiidae), rohu (Labeo rohita), rhinofishes (Labeo spp.), barramundi and giant sea perch (Lates calcarifer), striped mullet (Mugil cephalus), mullets (Mugilidae) (Mugil spp. and Liza spp.), ayu (Plecoglossus altivelis), pool barb (Puntius sophore), barcoo grunter (Scortum barcoo), sand whiting (Sillago ciliata), wells catfishes (Siluridae), snakeskin gourami (Trichogaster pectoralis), common archer fish (Toxotes chatareus), silver barb (Puntius gonionotus), spotted scat (Scatophagus argus), giant gourami (Osphronemus goramy), dusky flathead (Platycephalus fuscus), spiny turbot (Psettodes sp.), Tairiku-baratanago (Rhodeus ocellatus), Keti-Bangladeshi (Rohtee spp.), rudd (Scardinius erythrophthalmus), therapon (Terapon sp.) and three-spot gouramy (Trichogaster trichopterus). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.2.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with A. invadans status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with A. invadans, regardless of the infection with A. invadans status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animals and aquatic animal products from the species referred to in Article 10.2.2. which are intended for any purpose and which comply with Article 5.4.1.:

a) heat sterilised hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);

b) pasteurised fish products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent which has been demonstrated to inactivate A. invadans);

c) mechanically dried eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate A. invadans);

d) fish oil;

e) fish meal;

f) frozen eviscerated fish;

g) frozen fish fillets or steaks.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.2.2., other than those referred to in point 1 of Article 10.2.3., Competent Authorities should require the
conditions prescribed in Articles 10.2.7. to 10.2.11. relevant to infection with *A. invadans* status of the *exporting country, zone or compartment*.

3) When considering the importation or transit of *aquatic animals* and *aquatic animal products* from an *exporting country, zone or compartment* not declared free from infection with *A. invadans* of a species not covered in Article 10.2.2. but which could reasonably be expected to pose a *risk* of spread of infection with *A. invadans*, the *Competent Authority* should conduct a *risk analysis* in accordance with the recommendations in Chapter 2.1. The *exporting country* should be informed of the outcome of this assessment.

**Article 10.2.4.**

*Country free from infection with* *A. invadans*

If a country shares a *zone* with one or more other countries, it can only make a *self-declaration of freedom* from infection with *A. invadans* if all the areas covered by the shared water bodies are declared countries or *zones* free from infection with *A. invadans* (see Article 10.2.5.).

As described in Article 1.4.6., a country may make a *self-declaration of freedom* from infection with *A. invadans* if:

1) a country where there has been no observed occurrence of infection with *A. invadans* for at least the last ten years despite conditions that are conducive to its clinical expression, as described in the corresponding chapter of the *Aquatic Manual*, may make a *self-declaration of freedom* from infection with *A. invadans* when *basic biosecurity conditions* have been continuously met in the country for at least the last ten years;

OR

2) the disease status prior to *targeted surveillance* is unknown but the following conditions have been met:
   a) *basic biosecurity conditions* have been continuously met for at least the last two years; and
   b) *targeted surveillance*, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with *A. invadans*;

OR

3) it previously made a *self-declaration of freedom* from infection with *A. invadans* and subsequently lost its disease free status due to the detection of infection with *A. invadans* but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an *infected zone* and a *protection zone* was established; and
   b) infected populations have been destroyed or removed from the *infected zone* by means that minimise the risk of further spread of the disease, and the appropriate *disinfection* procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing *basic biosecurity conditions* have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) *targeted surveillance*, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with *A. invadans*.

In the meantime, part or all of the non-affected area may be declared a free *zone* provided that such a part meets the conditions in point 2 of Article 10.2.5.

**Article 10.2.5.**

*Zone or compartment free from infection with* *A. invadans*

If a *zone or compartment* extends over more than one country, it can only be declared a *zone or compartment* free from infection with *A. invadans* if all the relevant *Competent Authorities* confirm that all relevant conditions have been met.

As described in Article 1.4.6., a *zone or compartment* within the *territory* of one or more countries not declared free from infection with *A. invadans* may be declared free by the *Competent Authority(ies)* of the country(ies) concerned if:

1) a *zone or compartment* where the species referred to in Article 10.2.2. are present but there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression, as described in the corresponding chapter of the *Aquatic Manual*, may be declared free from infection
with *A. invadans* when basic biosecurity conditions have been continuously met in the zone or compartment for at least the last ten years;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with *A. invadans*;

OR

3) it previously made a self-declaration of freedom for a zone from infection with *A. invadans* and subsequently lost its disease free status due to the detection of infection with *A. invadans* in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with *A. invadans*.

Article 10.2.6.

**Maintenance of free status**

A country, zone or compartment that is declared free from infection with *A. invadans* following the provisions of point 1 of Articles 10.2.4. or 10.2.5. (as relevant) may maintain its status as free from infection with *A. invadans* provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with *A. invadans* following the provisions of point 2 of Articles 10.2.4. or 10.2.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with *A. invadans* provided that conditions that are conducive to clinical expression, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with *A. invadans*, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.2.7.

**Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with *A. invadans***

When importing aquatic animals and aquatic animal products of species referred to in Article 10.2.2. from a country, zone or compartment declared free from infection with *A. invadans*, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.2.4. or 10.2.5. (as applicable) and 10.2.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with *A. invadans*.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.2.3.
Chapter 10.2.- Infection with Aphanomyces invadans

Article 10.2.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with A. invadans

When importing for aquaculture, aquatic animals of a species referred to in Article 10.2.2. from a country, zone or compartment not declared free from infection with A. invadans, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive A. invadans in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with A. invadans.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for A. invadans in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with A. invadans (as described in Chapter 2.3.2. of the Aquatic Manual) and test for A. invadans in accordance with Chapter 1.4.;
      v) if A. invadans is not detected in the F-1 population, it may be defined as free from infection with A. invadans and may be released from quarantine;
      vi) if A. invadans is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.2.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with A. invadans

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.2.2. from a country, zone or compartment not declared free from infection with A. invadans, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.2.3., or products described in point 1 of Article 10.2.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of A. invadans or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 10.2.- Infection with Aphanomyces invadans

Article 10.2.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with A. invadans

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.2.2. from a country, zone or compartment not declared free from infection with A. invadans, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and
2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of A. invadans.

This Article does not apply to commodities referred to in point 1 of Article 10.2.3.

Article 10.2.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with A. invadans

1) Competent Authorities should not require any conditions related to infection with A. invadans, regardless of the infection with A. invadans status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (chilled) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.2.2. from a country, zone or compartment not declared free from infection with A. invadans, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.3.

INFECTION WITH GYRODACTYLUS SALARIS

Article 10.3.1.

For the purposes of the Aquatic Code, gyrodactylosis means infection with the viviparous freshwater ectoparasite Gyrodactylus salaris (Phylum Platyhelminthes; Class Monogenea).

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 10.3.2.

Scope

The recommendations in this chapter apply to: Atlantic salmon (Salmo salar), rainbow trout (Oncorhynchus mykiss), Arctic char (Salvelinus alpinus), North American brook trout (Salvelinus fontinalis), grayling (Thymallus thymallus), North American lake trout (Salvelinus namaycush) and brown trout (Salmo trutta). The recommendations also apply to other fish species in waters where the parasite is present, because these species may carry the parasite and act as vectors.

Article 10.3.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with G. salaris status of the exporting country, zone or compartment

1) Competent Authorities should not require any related conditions related to infection with G. salaris, regardless of the infection with G. salaris status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animals and aquatic animal products from the species referred to in Article 10.3.2. which are intended for any purpose and which comply with Article 5.4.1.:
   a) heat sterilised, hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) pasteurised fish products that have been subjected to a heat treatment at 63°C for at least 30 minutes (or any time/temperature equivalent which has been demonstrated to inactivate G. salaris);
   c) mechanically dried, eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate G. salaris);
   d) naturally dried, eviscerated fish (i.e. sun-dried or wind-dried);
   e) frozen eviscerated fish that have been subjected to minus 18°C or lower temperatures;
   f) frozen fish fillets or steaks that have been subjected to minus 18°C or lower temperatures;
   g) chilled eviscerated fish that have been harvested from seawater with a salinity of at least 25 parts per thousand (ppt);
   h) chilled fish fillets or steaks derived from fish that have been harvested from seawater with a salinity of at least 25 ppt;
   i) chilled fish products from which the skin, fins and gills have been removed;
   j) fish roe;
   k) fish oil;
   l) fish meal;
   m) fish skin leather.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.3.2., other than those referred to in point 1 of Article 10.3.3., Competent Authorities should require the conditions prescribed in Articles 10.3.7. to 10.3.11. relevant to the infection with G. salaris status of the exporting country, zone or compartment.
Chapter 10.3. - Infection with Gyrodactylus salaris

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 10.3.2. but which could reasonably be expected to pose a risk of spread of infection with G. salaris, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 10.3.4.

Country free from infection with G. salaris

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with G. salaris if all the areas covered by the shared watercourse(s) are declared countries or zones free from infection with G. salaris (see Article 10.3.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with G. salaris if:

1) none of the susceptible species referred to in Article 10.3.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.3.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last five years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last five years without detection of infection with G. salaris;

OR

4) it previously made a self-declaration of freedom from infection with G. salaris and subsequently lost its disease free status due to the detection of infection with G. salaris but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed, or the waters containing the infected fish have been treated by chemicals that kill the parasite; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last five years without detection of infection with G. salaris.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.3.5.

Article 10.3.5.

Zone or compartment free from infection with G. salaris

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with G. salaris if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with G. salaris may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 10.3.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;
2) any of the susceptible species referred to in Article 10.3.2. are present in the zone or compartment and the following conditions have been met:

   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and

   b) basic biosecurity conditions have been continuously met for at least the last five years;

OR

3) a zone or compartment supplied with seawater with a salinity of at least 25 ppt may be declared free from infection with G. salaris provided that no aquatic animals of species referred to in Article 10.3.2. are introduced from a site of a lesser health status for G. salaris during the 14 days prior to any live fish transfers from the zone or compartment;

OR

4) the disease status prior to targeted surveillance is unknown but the following conditions have been met:

   a) basic biosecurity conditions have been continuously met for at least the last ten years; and

   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last five years without detection of infection with G. salaris;

OR

5) it previously made a self-declaration of freedom for a zone from infection with G. salaris and subsequently lost its disease free status due to the detection of infection with G. salaris in the zone but the following conditions have been met:

   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and

   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed, or the waters containing the infected fish have been treated by chemicals that kill the parasite; and

   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and

   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last five years without detection of infection with G. salaris.

Article 10.3.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with G. salaris following the provisions of points 1 or 2 of Articles 10.3.4. or 10.3.5. (as relevant) may maintain its status as free from infection with G. salaris provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with G. salaris following the provisions of point 3 of Article 10.3.4. or point 4 of 10.3.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with G. salaris provided that conditions that are conducive to clinical expression of infection with G. salaris, as described in the corresponding chapter of the Aquatic Manual, exist, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with G. salaris, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.
Chapter 10.3. - Infection with Gyrodactylus salaris

Article 10.3.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with G. salaris

When importing aquatic animals and aquatic animal products of species referred to in Article 10.3.2. from a country, zone or compartment declared free from infection with G. salaris, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.3.4. or 10.3.5. (as applicable) and 10.3.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with G. salaris.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.3.3.

Article 10.3.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with G. salaris

When importing for aquaculture, aquatic animals of a species referred to in Article 10.3.2. from a country, zone or compartment not declared free from infection with G. salaris, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive G. salaris in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with G. salaris.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for G. salaris in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with G. salaris (as described in Chapter 2.3.3. of the Aquatic Manual) and test for G. salaris in accordance with Chapter 1.4.;
      v) if G. salaris is not detected in the F-1 population, it may be defined as free from infection with G. salaris and may be released from quarantine;
      vi) if G. salaris is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.
Chapter 10.3.- Infection with Gyrodactylus salaris

Article 10.3.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with G. salaris

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.3.2. from a country, zone or compartment not declared free from infection with G. salaris, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.3.3., or products described in point 1 of Article 10.3.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of G. salaris or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

Article 10.3.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with G. salaris

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.3.2. from a country, zone or compartment not declared free from infection with G. salaris, the Competent Authority of the importing country should:

1) require an international aquatic animal health certificate issued by the Competent Authority of the exporting country attesting that the aquatic animals have been held, immediately prior to export, in water with a salinity of at least 25 ppt for a continuous period of at least 14 days, and no other aquatic animals of the species referred to in Article 10.3.2. have been introduced during that period;

OR

2) require that the consignment be delivered directly to and held in quarantine facilities for slaughter and processing into one of the products referred to in point 1 of Article 10.3.3. or other products authorised by the Competent Authority, and water used in transport and all effluent and waste materials be treated in a manner that ensures inactivation of G. salaris.

This article does not apply to commodities referred to in point 1 of Article 10.3.3.

Article 10.3.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with G. salaris

1) Competent Authorities should not require any conditions related to infection with G. salaris, regardless of the infection with G. salaris status of the exporting country, zone or compartment, when authorising the importation or transit of the following commodities which have been prepared and packaged for retail trade and which comply with Article 5.4.2.:

   – no commodities listed.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.3.2. from a country, zone or compartment not declared free from infection with G. salaris, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 1997; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.4.

INFECTION WITH INFECTIOUS SALMON ANAEMIA VIRUS

Article 10.4.1.

For the purposes of the Aquatic Code, infection with infectious salmon anaemia virus (ISAV) means infection with HPR0 (non-deleted highly polymorphic region) or HPR-deleted ISAV of the genus Isavirus of the family Orthomyxoviridae. Both genotypes should be notified in accordance with the Aquatic Code.

There is a link between non-pathogenic HPR0 ISAV and pathogenic HPR-deleted ISAV, with some outbreaks potentially occurring as a result of the emergence of HPR-deleted from HPR0.

The provisions in this chapter are provided in recognition of three possible levels of disease status with respect to ISAV:
1) HPR0 ISAV and HPR-deleted ISAV free;
2) HPR0 ISAV endemic (but HPR-deleted ISAV free);
3) HPR0 ISAV and HPR-deleted ISAV endemic.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 10.4.2.

Scope

The recommendations in this chapter apply to: Atlantic salmon (Salmo salar), brown trout (Salmo trutta) and rainbow trout (Onchorynchus mykiss). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.4.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infectious salmon anaemia virus status of the exporting country, zone or compartment

In this article, all statements referring to ISAV are for any detectable ISAV, including HPR0 ISAV.

1) Competent Authorities should not require any conditions related to infection with ISAV, regardless of the infection with ISAV status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.4.2, which are intended for any purpose and which comply with Article 5.4.1.:
   a) heat sterilised, hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) pasteurised fish products that have been subjected to a heat treatment at 90°C for at least ten minutes (or to any time/temperature equivalent which has been demonstrated to inactivate ISAV);
   c) mechanically dried, eviscerated fish (i.e. a heat treatment at 100°C for 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate ISAV);
   d) fish oil;
   e) fish meal;
   f) fish skin leather.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.4.2., other than those referred to in point 1 of Article 10.4.3., Competent Authorities should require the conditions prescribed in Articles 10.4.10. to 10.4.16. relevant to the ISAV status of the exporting country, zone or compartment.
3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 10.4.2, but which could reasonably be expected to pose a risk of spread of infection with ISAV, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 10.4.4.

Country free from infection with infectious salmon anaemia virus

In this article, all statements referring to a country free from infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with ISAV if all the areas covered by the shared water bodies are declared countries or zones free from infection with ISAV (see Article 10.4.6.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with ISAV if:

1) none of the susceptible species referred to in Article 10.4.2 are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with ISAV;

OR

3) it previously made a self-declaration of freedom from infection with ISAV and subsequently lost its disease free status due to the detection of infection with ISAV but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with ISAV.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.4.6.

The pathway for self-declaration of freedom from infection with ISAV HPR0 based on absence of clinical disease (referred to as historical freedom in Article 1.4.6.) cannot be achieved because infection with ISAV HPR0 is unlikely to cause any clinical signs.

Article 10.4.5.

Country free from infection with HPR-deleted infectious salmon anaemia virus

In this article, all statements refer to a country free from infection with HPR-deleted ISAV but not necessarily free from infection with HPR0 ISAV.

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with HPR-deleted ISAV if all the areas covered by the shared water bodies are declared countries or zones free from infection with HPR-deleted ISAV (see Article 10.4.7.).
Chapter 10.4.- Infection with infectious salmon anaemia virus

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with HPR-deleted ISAV if:

1) any of the susceptible species referred to in Article 10.4.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of infection with HPR-deleted ISAV for at least the last ten years despite conditions that are conducive to clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with HPR-deleted ISAV;

OR

3) it previously made a self-declaration of freedom from infection with HPR-deleted ISAV and subsequently lost its disease free status due to the detection of infection with HPR-deleted ISAV but the following conditions have been met:
   a) on detection of infection with HPR-deleted ISAV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of HPR-deleted ISAV, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with HPR-deleted ISAV.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.4.7.

Article 10.4.6.

Zone or compartment free from infection with infectious salmon anaemia virus

In this article, all statements referring to a zone or compartment free from infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with ISAV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with ISAV may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 10.4.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with ISAV;
3) it previously made a self-declaration of freedom for a zone from infection with ISAV and subsequently lost its disease free status due to the detection of infection with ISAV in the zone but the following conditions have been met:
   a) on detection of infection with ISAV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with ISAV.

Article 10.4.7.

Zone or compartment free from infection with HPR-deleted infectious salmon anaemia virus

In this article, all statements refer to a zone or compartment free from infection with HPR-deleted ISAV but not necessarily free from infection with HPR0 ISAV.

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with HPR-deleted ISAV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with HPR-deleted ISAV may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) any of the susceptible species referred to in Article 10.4.2. are present in the zone or compartment and the following conditions have been met:
   a) there has been no observed occurrence of infection with HPR-deleted ISAV for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with HPR-deleted ISAV;

OR

3) it previously made a self-declaration of freedom for a zone from infection with HPR-deleted ISAV and subsequently lost its disease free status due to the detection of infection with HPR-deleted ISAV in the zone but the following conditions have been met:
   a) on detection of infection with HPR-deleted ISAV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least two years without detection of infection with HPR-deleted ISAV.
Article 10.4.8.

Maintenance of free status for infection with infectious salmon anaemia virus

In this article, all statements referring to a country, zone or compartment free from infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

A country, zone or compartment that is declared free from infection with ISAV following the provisions of point 1 of Articles 10.4.4. or 10.4.6. (as relevant) may maintain its status as free from infection with ISAV provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with ISAV following the provisions of point 2 of Articles 10.4.4. or 10.4.6. (as relevant) may maintain its status as free from infection with ISAV provided that targeted surveillance is continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection, and basic biosecurity conditions are continuously maintained.

Article 10.4.9.

Maintenance of free status for infection with HPR-deleted infectious salmon anaemia virus

In this article, all statements refer to a country, zone or compartment free from infection with HPR-deleted ISAV, but not necessarily free from infection with HPR0 ISAV.

A country, zone or compartment that is declared free from infection with HPR-deleted ISAV following the provisions of points 1 or 2 of Articles 10.4.5. or 10.4.7. (as relevant) may maintain its status as free from infection with ISAV provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with HPR-deleted ISAV following the provisions of point 3 of Articles 10.4.5. or 10.4.7. (as relevant) may discontinue targeted surveillance and maintain its free status provided that conditions that are conducive to clinical expression, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in an infected country and in all cases where conditions are not conducive to clinical expression, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.4.10.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with infectious salmon anaemia virus

In this article, all statements referring to a country, zone or compartment free from infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

When importing aquatic animals and aquatic animal products of species referred to in Article 10.4.2. from a country, zone or compartment declared free from infection with ISAV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.4.4. or 10.4.6. (as applicable) and 10.4.8., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with ISAV.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.4.3.
Chapter 10.4.- Infection with infectious salmon anaemia virus

Article 10.4.11.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with HPR-deleted infectious salmon anaemia virus

In this article, all statements refer to a country, zone or compartment free from infection with HPR-deleted ISAV, but not necessarily free from infection with HPR0 ISAV.

When importing aquatic animals and aquatic animal products of species referred to in Article 10.4.2. from a country, zone or compartment declared free from infection with HPR-deleted ISAV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.4.5. or 10.4.7. (as applicable) and 10.4.9., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with HPR-deleted ISAV.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.4.3.

Article 10.4.12.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with infectious salmon anaemia virus

In this article, all statements referring to infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

When importing for aquaculture, aquatic animals of a species referred to in Article 10.4.2. from a country, zone or compartment not declared free from infection with ISAV, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive ISAV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with ISAV.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for ISAV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with ISAV (as described in Chapter 2.3.5. of the Aquatic Manual) and test for ISAV in accordance with Chapter 1.4.;
      v) if ISAV is not detected in the F-1 population, it may be defined as free from infection with ISAV and may be released from quarantine;
      vi) if ISAV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.
Chapter 10.4.- Infection with infectious salmon anaemia virus

Article 10.4.13.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with infectious salmon anaemia virus

In this article, all statements referring to infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.4.2. from a country, zone or compartment not declared free from infection with ISAV, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.4.3., or products described in point 1 of Article 10.4.15., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of ISAV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

Article 10.4.14.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with infectious salmon anaemia virus

In this article, all statements referring to infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.4.2. from a country, zone or compartment not declared free from infection with ISAV, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of ISAV.

This article does not apply to commodities referred to in point 1 of Article 10.4.3.

Article 10.4.15.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with infectious salmon anaemia virus

In this article, all statements referring to infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

1) Competent Authorities should not require any conditions related to infection with ISAV, regardless of the infection with ISAV status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (frozen or chilled) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.4.2. from a country, zone or compartment not declared free from infection with ISAV, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.
Chapter 10.4.- Infection with infectious salmon anaemia virus

Article 10.4.16.

Importation of disinfected eggs for aquaculture from a country, zone or compartment not declared free from infection with infectious salmon anaemia virus

In this article, all statements referring to infection with ISAV are for any detectable ISAV, including HPR0 ISAV.

1) When importing disinfected eggs of the species referred to in Article 10.4.2. for aquaculture, from a country, zone or compartment not declared free from infection with ISAV, the Competent Authority of the importing country should assess the risk associated with at least:
   a) the ISAV status of the water to be used during the disinfection of the eggs;
   b) the level of infection with ISAV in broodstock (ovarian fluid and milt); and
   c) the temperature and pH of the water to be used for disinfection.

2) If the Competent Authority of the importing country concludes that the importation is acceptable, it should apply the following risk mitigation measures including:
   a) the eggs should be disinfected prior to importing, in accordance with recommendations in Chapter 4.4. or those specified by the Competent Authority of the importing country; and
   b) between disinfection and the import, eggs should not come into contact with anything which may affect their health status.

The Competent Authority may wish to consider internal measures, such as renewed disinfection of the eggs upon arrival in the importing country.

3) When importing disinfected eggs of the species referred to in Article 10.4.2. for aquaculture, from a country, zone or compartment not declared free from infection with ISAV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that the procedures described in point 2 of this article have been fulfilled.

NB: FIRST ADOPTED IN 1995; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.5.

INFECTION WITH SALMONID ALPHAVIRUS

Article 10.5.1.

General provisions

For the purposes of the Aquatic Code, infection with salmonid alphavirus (SAV) means infection with any subtype of SAV of the genus Alphavirus of the family Togaviridae.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 10.5.2.

Scope

The recommendations in this chapter apply to: Atlantic salmon (Salmo salar), brown trout (Salmo trutta) and rainbow trout (Onchorynchus mykiss). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.5.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with salmonid alphavirus status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with SAV, regardless of the infection with SAV status of the exporting country, zone or compartment when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.5.2. intended for any purpose and complying with Article 5.4.1.:
   a) heat sterilised, hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) pasteurised fish products that have been subjected to a heat treatment at 90°C for at least ten minutes (or to any time/temperature equivalent which has been demonstrated to inactivate SAV);
   c) mechanically dried, eviscerated fish (i.e. a heat treatment at 100°C for 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate SAV);
   d) fish oil;
   e) fish meal;
   f) fish skin leather.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.5.2., other than those referred to in point 1 of Article 10.5.3., Competent Authorities should require the conditions prescribed in Articles 10.5.7. to 10.5.12. relevant to the SAV status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 10.5.2. but which could reasonably be expected to pose a risk of spread of infection with SAV, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.
Chapter 10.5. - Infection with salmonid alphavirus

Article 10.5.4.

Country free from infection with salmonid alphavirus

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with SAV if all the areas covered by the shared water bodies are declared countries or zones free from infection with SAV (see Article 10.5.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with SAV if:

1) none of the susceptible species referred to in Article 10.5.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.5.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the past ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with SAV;

OR

4) it previously made a self-declaration of freedom from infection with SAV and subsequently lost its disease free status due to the detection of infection with SAV but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with SAV.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.5.5.

Article 10.5.5.

Zone or compartment free from infection with salmonid alphavirus

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with SAV if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with SAV may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 10.5.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.5.2. are present in the zone or compartment and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;
3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with SAV;

OR

4) it previously made a self-declaration of freedom for a zone from infection with SAV and subsequently lost its disease free status due to the detection of infection with SAV in the zone but the following conditions have been met:
   a) on detection of infection with SAV, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with SAV.

Article 10.5.6.

Maintenance of free status for infection with salmonid alphavirus

A country, zone or compartment that is declared free from infection with SAV following the provisions of points 1 or 2 of Articles 10.5.4. or 10.5.5. (as relevant) may maintain its status as free from infection with SAV provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with SAV following the provisions of point 3 of Articles 10.5.4. or 10.5.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with SAV provided that conditions that are conducive to clinical expression, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in an infected country and in all cases where conditions are not conducive to clinical expression, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.5.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with salmonid alphavirus

When importing aquatic animals and aquatic animal products of species referred to in Article 10.5.2. from a country, zone or compartment declared free from infection with SAV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.5.4. or 10.5.5. (as applicable) and 10.5.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with SAV.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.5.3.
Article 10.5.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with salmonid alphavirus

When importing for aquaculture, aquatic animals of a species referred to in Article 10.5.2. from a country, zone or compartment not declared free from infection with SAV, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive SAV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with SAV.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for SAV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with SAV (as described in Chapter 2.3.6. of the Aquatic Manual) and test for SAV in accordance with Chapter 1.4.;
      v) if SAV is not detected in the F-1 population, it may be defined as free from infection with SAV and may be released from quarantine;
      vi) if SAV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.5.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with salmonid alphavirus

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.5.2. from a country, zone or compartment not declared free from infection with SAV, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.5.3., or products described in point 1 of Article 10.5.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of SAV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 10.5.- Infection with salmonid alphavirus

Article 10.5.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with salmonid alphavirus

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.5.2. from a country, zone or compartment not declared free from infection with SAV, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine for slaughter and processing into products authorised by the Competent Authority, and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of SAV.

This article does not apply to commodities referred to in point 1 of Article 10.5.3.

Article 10.5.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with salmonid alphavirus

1) Competent Authorities should not require any conditions related to infection with SAV, regardless of the infection with SAV status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (frozen or chilled) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.5.2. from a country, zone or compartment not declared free from infection with SAV, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

Article 10.5.12.

Importation of disinfected eggs for aquaculture from a country, zone or compartment not declared free from infection from infection with salmonid alphavirus

1) When importing disinfected eggs of the species referred to in Article 10.5.2. for aquaculture, from a country, zone or compartment not declared free from infection with SAV, the Competent Authority of the importing country should assess the risk associated with at least:

a) the SAV status of the water to be used during the disinfection of the eggs;

b) the level of infection with SAV in broodstock; and

c) the temperature and pH of the water to be used for disinfection.

2) If the Competent Authority of the importing country concludes that the importation is acceptable, it should apply the following risk mitigation measures including:

a) the eggs should be disinfected prior to importing, in accordance with recommendations in Chapter 4.4. or those specified by the Competent Authority of the importing country; and

b) between disinfection and the import, eggs should not come into contact with anything which may affect their health status.

The Competent Authority may wish to consider internal measures, such as renewed disinfection of the eggs upon arrival in the importing country.
3) When importing disinfected eggs of the species referred to in Article 10.5.2. for aquaculture, from a country, zone or compartment not declared free from infection with SAV, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that the procedures described in point 2 of this article have been fulfilled.

NB: FIRST ADOPTED IN 2014; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.6.

INFECTIOUS HAEMATOPOIETIC NECROSIS

Article 10.6.1.

For the purposes of the Aquatic Code, infectious haematopoietic necrosis (IHN) means infection with IHN virus (IHNV) of the genus Novirhabdovirus of the family Rhabdoviridae.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 10.6.2.

Scope

The recommendations in this chapter apply to: rainbow trout or steelhead (Oncorhynchus mykiss), the Pacific salmon species (chinook [Oncorhynchus tshawytscha], sockeye [Oncorhynchus nerka], chum [Oncorhynchus keta], masou [Oncorhynchus masou], pink [Oncorhynchus rhodurus] and coho [Oncorhynchus kisutch]), and Atlantic salmon (Salmo salar). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.6.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infectious haematopoietic necrosis status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to IHN, regardless of the IHN status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.6.2. which are intended for any purpose and which comply with Article 5.4.1.:
   a) heat sterilised, hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) pasteurised fish products that have been subjected to a heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent which has been demonstrated to inactivate IHNV);
   c) mechanically dried, eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate IHNV);
   d) fish oil;
   e) fish meal;
   f) fish skin leather.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.6.2., other than those referred to in point 1 of Article 10.6.3., Competent Authorities should require the conditions prescribed in Articles 10.6.7. to 10.6.12. relevant to the IHN status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 10.6.2. but which could reasonably be expected to pose a risk of spread of IHN, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.
Article 10.6.4.

Country free from infectious haematopoietic necrosis

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from IHN if all the areas covered by the shared water bodies are declared countries or zones free from IHN (see Article 10.6.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from IHN if:

1) none of the susceptible species referred to in Article 10.6.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.6.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IHN;

OR

4) it previously made a self-declaration of freedom from IHN and subsequently lost its disease free status due to the detection of IHN but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IHN.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.6.5.

Article 10.6.5.

Zone or compartment free from infectious haematopoietic necrosis

If a zone or compartment extends over more than one country, it can only be declared an IHN free zone or compartment if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from IHN may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 10.6.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.6.2. are present in the zone or compartment and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;
3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of IHN;

OR

4) it previously made a self-declaration of freedom for a zone from IHN and subsequently lost its disease free status due to the detection of IHN in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of IHN.

Article 10.6.6.

Maintenance of free status

A country, zone or compartment that is declared free from IHN following the provisions of points 1 or 2 of Articles 10.6.4. or 10.6.5. (as relevant) may maintain its status as free from IHN provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from IHN following the provisions of point 3 of Articles 10.6.4. or 10.6.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from IHN provided that conditions that are conducive to clinical expression of IHN, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of IHN, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.6.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infectious haematopoietic necrosis

When importing aquatic animals and aquatic animal products of species referred to in Article 10.6.2 from a country, zone or compartment declared free from IHN, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.6.4. or 10.6.5. (as applicable) and 10.6.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from IHN.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.6.3.
Chapter 10.6.- Infectious haematopoietic necrosis

Article 10.6.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infectious haematopoietic necrosis

When importing for aquaculture, aquatic animals of a species referred to in Article 10.6.2. from a country, zone or compartment not declared free from IHN, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive IHNV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for IHN.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for IHNV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of IHN (as described in Chapter 2.3.4. of the Aquatic Manual) and test for IHNV in accordance with Chapter 1.4.;
      v) if IHNV is not detected in the F-1 population, it may be defined as free from IHN and may be released from quarantine;
      vi) if IHNV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.6.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infectious haematopoietic necrosis

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.6.2. from a country, zone or compartment not declared free from IHN, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.6.3., or products described in point 1 of Article 10.6.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of IHNV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 10.6.- Infectious haematopoietic necrosis

Article 10.6.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infectious haematopoietic necrosis

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.6.2. from a country, zone or compartment not declared free from IHN, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of IHNV.

This article does not apply to commodities referred to in point 1 of Article 10.6.3.

Article 10.6.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infectious haematopoietic necrosis

1) Competent Authorities should not require any conditions related to IHN, regardless of the IHN status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (frozen or chilled) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.6.2. from a country, zone or compartment not declared free from IHN, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

Article 10.6.12.

Importation of disinfected eggs for aquaculture from a country, zone or compartment not declared free from infectious haematopoietic necrosis

1) When importing disinfected eggs of the species referred to in Article 10.6.2. for aquaculture, from a country, zone or compartment not declared free from IHN, the Competent Authority of the importing country should assess the risk associated with at least:

   a) the IHN virus status of the water to be used during the disinfection of the eggs;

   b) the prevalence of infection with IHN virus in broodstock (ovarian fluid and milt); and

   c) the temperature and pH of the water to be used for disinfection.

2) If the Competent Authority of the importing country concludes that the importation is acceptable, it should apply the following risk mitigation measures including:

   a) the eggs should be disinfected prior to importing, in accordance with recommendations in Chapter 4.4. or those specified by the Competent Authority of the importing country; and

   b) between disinfection and the import, eggs should not come into contact with anything which may affect their health status.

The Competent Authority may wish to consider internal measures, such as renewed disinfection of the eggs upon arrival in the importing country.
3) When importing disinfected eggs of species referred to in Article 10.6.2. for aquaculture, from a country, zone or compartment not declared free from IHN, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that the procedures described in point 2 of Article 10.6.12. have been fulfilled.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.7.

KOI HERPESVIRUS DISEASE

Article 10.7.1.

For the purposes of the Aquatic Code, koi herpesvirus disease (KHVD) means infection with the viral species koi herpesvirus (KHV) tentatively placed in the sub-family Cyprinid herpesvirus of the family Herpesviridae.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 10.7.2.

Scope

The recommendations in this chapter apply to: common carp (Cyprinus carpio carpio), ghost carp (Cyprinus carpio goi), koi carp (Cyprinus carpio koi) and common carp hybrids (e.g. Cyprinus carpio xCarassius auratus). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.7.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the koi herpesvirus disease status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to KHVD, regardless of the KHVD status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.7.2. which are intended for any purpose and which comply with Article 5.4.1.:
   a) heat sterilised hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) pasteurised fish products that have been subjected to heat treatment at 90°C for at least ten minutes (or to any time/temperature equivalent which has been demonstrated to inactivate KHV);
   c) mechanically dried eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes (or any time/temperature equivalent which has been demonstrated to inactivate KHV);
   d) fish oil;
   e) fish meal.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.7.2., other than those referred to in point 1 of Article 10.7.3., Competent Authorities should require the conditions prescribed in Articles 10.7.7. to 10.7.11. relevant to the KHVD status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 10.7.2. but which could reasonably be expected to pose a risk of spread of KHVD, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 10.7.4.

Country free from koi herpesvirus disease

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from KHVD if all the areas covered by the shared water bodies are declared countries or zones free from KHVD (see Article 10.7.5.).
Chapter 10.7.- Koi herpesvirus disease

As described in Article 1.4.6., a country may make a self-declaration of freedom from KHVD if:

1) none of the susceptible species referred to in Article 10.7.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.7.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of KHVD;

OR

4) it previously made a self-declaration of freedom from KHVD and subsequently lost its disease free status due to the detection of KHVD but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of KHVD.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.7.5.

Article 10.7.5.

Zone or compartment free from koi herpesvirus disease

If a zone or compartment extends over more than one country, it can only be declared a KHVD free zone or compartment if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from KHVD may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 10.7.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.7.2. are present in the zone or compartment and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of KHVD.
OR

4) it previously made a self-declaration of freedom for a zone from KHVD and subsequently lost its disease free status due to the detection of KHVD in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of KHVD.

Article 10.7.6.

Maintenance of free status

A country, zone or compartment that is declared free from KHVD following the provisions of points 1 or 2 of Articles 10.7.4. or 10.7.5. (as relevant) may maintain its status as free from KHVD provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from KHVD following the provisions of point 3 of Articles 10.7.4. or 10.7.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from KHVD provided that conditions that are conducive to clinical expression of KHVD, as described in the corresponding chapter of the Aquatic Manual, exist, and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of KHVD, targeted surveillance needs to be continued at a level determined by the Competent Authority on the basis of the likelihood of infection.

Article 10.7.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from koi herpesvirus disease

When importing aquatic animals and aquatic animal products of species referred to in Article 10.7.2. from a country, zone or compartment declared free from KHVD, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.7.4. or 10.7.5. (as applicable) and 10.7.6., the place of production of aquatic animals and aquatic animal products is a country, zone or compartment declared free from KHVD.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.7.3.

Article 10.7.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from koi herpesvirus disease

When importing for aquaculture, aquatic animals of a species referred to in Article 10.7.2. from a country, zone or compartment not declared free from KHVD, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive KHV in accordance with Chapters 4.3., 4.7. and 5.5.
2) If the intention is to establish a new stock for aquaculture, consider applying the following:

a) In the exporting country:
   i) identify potential source populations and evaluate their aquatic animal health records;
   ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for KHVD.

b) In the importing country:
   i) import the F-0 population into a quarantine facility;
   ii) test the F-0 population for KHV in accordance with Chapter 1.4. to determine their suitability as broodstock;
   iii) produce a first generation (F-1) population in quarantine;
   iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of KHVD (as described in Chapter 2.3.7. of the Aquatic Manual) and test for KHV in accordance with Chapter 1.4.;
   v) if KHV is not detected in the F-1 population, it may be defined as free from KHVD and may be released from quarantine;
   vi) if KHV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.7.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from koi herpesvirus disease

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.7.2. from a country, zone or compartment not declared free from KHVD, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.7.3., or products described in point 1 of Article 10.7.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of KHV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

Article 10.7.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from koi herpesvirus disease

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of a species referred to in Article 10.7.2. from a country, zone or compartment not declared free from KHVD, the Competent Authority of the importing country should require that:

1) the consignment be delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing be treated in a manner that ensures inactivation of KHV.

This article does not apply to commodities referred to in point 1 of Article 10.7.3.
Article 10.7.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from koi herpesvirus disease

1) Competent Authorities should not require any conditions related to KHVD, regardless of the KHVD status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (frozen or chilled) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.7.2. from a country, zone or compartment not declared free from KHVD, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2007; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.8.

RED SEA BREAM IRIDO VIRAL DISEASE

Article 10.8.1.

For the purposes of the Aquatic Code, red sea bream iridoviral disease (RSIVD) means infection with red sea bream iridovirus (RSIV) of the family Iridoviridae.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 10.8.2.

Scope

The recommendations in this chapter apply to: red sea bream (*Pagrus major*), yellowtail (*Seriola quinqueradiata*), amberjack (*Seriola dumerili*), sea bass (*Lateolabrax sp.* and *Lates calcarifer*), Albacore (*Thunnus thynnus*), Japanese parrotfish (*Oplegnathus fasciatus*), striped jack (*Caranx delicatissimus*), mandarin fish (*Siniperca chuatsi*), red drum (*Sciaenops ocellatus*), mullet (*Mugil cephalus*) and groupers (*Epinephelus* spp.). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.8.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the red sea bream iridoviral disease status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to RSIVD, regardless of the RSIVD status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.8.2. which are intended for any purpose and which comply with Article 5.4.1.:

   a) heat sterilised hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   
   b) pasteurised fish products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent which has been demonstrated to inactivate RSIV);
   
   c) mechanically dried eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes (or any time/temperature equivalent which has been demonstrated to inactivate RSIV);
   
   d) fish oil;
   
   e) fish meal;
   
   f) fish skin leather.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.8.2., other than those referred to in point 1 of Article 10.8.3., Competent Authorities should require the conditions prescribed in Articles 10.8.7. to 10.8.11. relevant to the RSIVD status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 10.8.2. but which could reasonably be expected to pose a risk of spread of RSIVD, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.
Chapter 10.8. - Red sea bream iridoviral disease

Article 10.8.4.

Red sea bream iridovirus free country

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from RSIVD if all the areas covered by the shared water bodies are declared countries or zones free from RSIVD (see Article 10.8.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from RSIVD if:

1) none of the susceptible species referred to in Article 10.8.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.8.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual), and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of RSIV;

OR

4) it previously made a self-declaration of freedom from RSIVD and subsequently lost its disease free status due to the detection of RSIV but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of RSIV.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 10.8.5.

Article 10.8.5.

Red sea bream iridoviral diseases free zone or free compartment

If a zone or compartment extends over more than one country, it can only be declared a RSIVD free zone or compartment if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from RSIVD may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 10.8.2. are present in the zone or compartment and basic biosecurity conditions have been met continuously for at least the last two years;

OR

2) any of the susceptible species referred to in Article 10.8.2. are present in the zone or compartment and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last ten years;
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OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of RSIV;

OR

4) it previously made a self-declaration of freedom for a zone from RSIVD and subsequently lost its disease free status due to the detection of RSIVD in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of RSIV.

Article 10.8.6.

Maintenance of free status

A country, zone or compartment that is declared free from RSIVD following the provisions of points 1 or 2 of Articles 10.8.4. or 10.8.5. (as relevant) may maintain its status as free from RSIVD provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from RSIVD following the provisions of point 3 of Articles 10.8.4. or 10.8.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from RSIVD provided that conditions that are conducive to clinical expression of RSIVD, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of RSIVD, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.8.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from red sea bream iridoviral disease

When importing aquatic animals and aquatic animal products of species referred to in Article 10.8.2. from a country, zone or compartment declared free from RSIVD, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.8.4. or 10.8.5. (as applicable) and 10.8.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from RSIVD.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.8.3.
Chapter 10.8. - Red sea bream iridoviral disease

Article 10.8.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from red sea bream iridoviral disease

When importing for aquaculture, aquatic animals of a species referred to in Article 10.8.2. from a country, zone or compartment not declared free from RSIVD, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive RSIV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for RSIVD.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for RSIV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of RSIVD (as described in Chapter 2.3.8. of the Aquatic Manual) and test for RSIV in accordance with Chapter 1.4.;
      v) if RSIV is not detected in the F-1 population, it may be defined as free from RSIVD and may be released from quarantine;
      vi) if RSIV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.8.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from red sea bream iridoviral disease

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.8.2. from a country, zone or compartment not declared free from RSIVD, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to, and held in, quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.8.3., or products described in point 1 of Article 10.8.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of RSIV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 10.8.- Red sea bream iridoviral disease

Article 10.8.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use from a country, zone or compartment not declared free from red sea bream iridoviral disease

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.8.2. from a country, zone or compartment not declared free from RSIVD, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of RSIV.

This article does not apply to commodities referred to in point 1 of Article 10.8.3.

Article 10.8.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from red sea bream iridoviral disease

1) Competent Authorities should not require any RSIVD related conditions, regardless of the RSIVD status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (chilled or frozen) which have been prepared and packaged for retail trade and which comply with Article 5.4.2. Certain assumptions have been made in assessing the safety of the aquatic animal products listed above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.8.2. from a country, zone or compartment not declared free from RSIVD, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.9.

SPRING VIRAEMIA OF CARP

Article 10.9.1.

For the purposes of the Aquatic Code, spring viraemia of carp (SVC) means infection with the viral species SVC virus (SVCV) tentatively placed in the genus Vesiculovirus of the family Rhabdoviridae.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 10.9.2.

Scope

The recommendations in this chapter apply to: common carp (Cyprinus carpio carpio) and koi carp (Cyprinus carpio koi), crucian carp (Carassius carassius), sheatfish (also known as European catfish or wels) (Silurus glanis), silver carp (Hypophthalmichthys molitrix), bighead carp (Aristichthys nobilis), grass carp (white amur) (Ctenopharyngodon idellus), goldfish (Carassius auratus), orfe (Leuciscus idus), and tench (Tinca tinca). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.9.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the spring viraemia of carp status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to SVC, regardless of the SVC status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.9.2. which are intended for any purpose and which comply with Article 5.4.1.:

   a) heat sterilised hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or equivalent);
   b) pasteurised fish products that have been subjected to heat treatment at 90°C for at least ten minutes (or any time/temperature equivalent which has been demonstrated to inactivate SVCV);
   c) mechanically dried eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate SVCV);
   d) fish oil;
   e) fish meal.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 10.9.2., other than those referred to in point 1 of Article 10.9.3., Competent Authorities should require the conditions prescribed in Articles 10.9.7. to 10.9.11. relevant to the SVC status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 10.9.2. but which could reasonably be expected to pose a risk of spread of SVC, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.
Chapter 10.9.- Spring viraemia of carp

Article 10.9.4.

Country free from spring viraemia of carp

If a country shares a zone with one or more other countries, it can only make a *self-declaration of freedom* from SVC if all the areas covered by the shared water bodies are declared countries or zones free from SVC (see Article 10.9.5).

As described in Article 1.4.6., a country may make a *self-declaration of freedom* from SVC if:

1) none of the *susceptible species* referred to in Article 10.9.2. are present and *basic biosecurity conditions* have been continuously met for at least the last two years;

OR

2) any of the *susceptible species* referred to in Article 10.9.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the *disease* for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) *basic biosecurity conditions* have been continuously met for at least the last ten years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) *basic biosecurity conditions* have been continuously met for at least the last two years; and
   b) *targeted surveillance*, as described in Chapter 1.4., has been in place for at least the last two years without detection of SVC;

OR

4) it previously made a *self-declaration of freedom* from SVC and subsequently lost its disease free status due to the detection of SVC but the following conditions have been met:
   a) on detection of the *disease*, the affected area was declared an *infected zone* and a *protection zone* was established; and
   b) infected populations have been destroyed or removed from the *infected zone* by means that minimise the risk of further spread of the disease, and the appropriate *disinfection* procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing *basic biosecurity conditions* have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) *targeted surveillance*, as described in Chapter 1.4., has been in place for at least the last two years without detection of SVC.

In the meantime, part or all of the non-affected area may be declared a free *zone* provided that such a part meets the conditions in point 3 of Article 10.9.5.

Article 10.9.5.

Zone or compartment free from spring viraemia of carp

If a *zone* or *compartment* extends over more than one country, it can only be declared an SVC free *zone* or *compartment* if all the relevant *Competent Authorities* confirm that all relevant conditions have been met.

As described in Article 1.4.6., a *zone* or *compartment* within the territory of one or more countries not declared free from SVC may be declared free by the *Competent Authority(ies)* of the country(ies) concerned if:

1) none of the *susceptible species* referred to in Article 10.9.2. are present in the *zone* or *compartment* and *basic biosecurity conditions* have been continuously met for at least the last two years;

OR

2) any of the *susceptible species* referred to in Article 10.9.2. are present in the *zone* or *compartment* and the following conditions have been met:
   a) there has been no observed occurrence of the *disease* for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) *basic biosecurity conditions* have been continuously met for at least the last ten years;
3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of SVC;

OR

4) it previously made a self-declaration of freedom for a zone from SVC and subsequently lost its disease free status due to the detection of SVC in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of SVC.

Article 10.9.6.

Maintenance of free status

A country, zone or compartment that is declared free from SVC following the provisions of points 1 or 2 of Articles 10.9.4. or 10.9.5. (as relevant) may maintain its status as free from SVC provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from SVC following the provisions of point 3 of Articles 10.9.4. or 10.9.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from SVC provided that conditions that are conducive to clinical expression of SVC, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of SVC, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.9.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from spring viraemia of carp

When importing aquatic animals and aquatic animal products of species referred to in Article 10.9.2. from a country, zone or compartment declared free from SVC, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.9.4. or 10.9.5. (as applicable) and 10.9.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from SVC.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.9.3.
Article 10.9.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from spring viraemia of carp

When importing for aquaculture, aquatic animals of a species referred to in Article 10.9.2. from a country, zone or compartment not declared free from SVC, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive SVCV in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for SVC.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for SVCV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of SVC (as described in Chapter 2.3.9. of the Aquatic Manual) and test for SVCV in accordance with Chapter 1.4.;
      v) if SVCV is not detected in the F-1 population, it may be defined as free from SVC and may be released from quarantine;
      vi) if SVCV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.9.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from spring viraemia of carp

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.9.2. from a country, zone or compartment not declared free from SVC, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.9.3., or products described in point 1 of Article 10.9.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of SVCV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Article 10.9.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use from a country, zone or compartment not declared free from spring viraemia of carp

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.9.2. from a country, zone or compartment not declared free from SVC, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of SVCV.

This article does not apply to commodities referred to in point 1 of Article 10.9.3.

Article 10.9.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from spring viraemia of carp

1) Competent Authorities should not require any conditions related to SVC, regardless of the SVC status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (chilled or frozen) which have been prepared and packaged for retail trade and which comply with Article 5.4.2. Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.9.2. from a country, zone or compartment not declared free from SVC, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 10.10.

VIRAL HAEMORRHAGIC SEPTICAEMIA

Article 10.10.1.

For the purposes of the Aquatic Code, viral haemorrhagic septicaemia (VHS) means infection with VHS virus (VHSV, synonym: Egtved virus) of the genus Novirhabdovirus of the family Rhabdoviridae.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 10.10.2.

Scope

The recommendations in this chapter apply to: rainbow trout (Oncorhynchus mykiss), brown trout (Salmo trutta), grayling (Thymallus thymallus), white fish (Coregonus spp.), pike (Esox lucius), turbot (Scophthalmus maximus), herring and sprat (Clupea spp.), Pacific salmon (Oncorhynchus spp.), Atlantic cod (Gadus morhua), Pacific cod (Gadus macrocephalus), haddock (Gadus aeglefinus) and rockling (Onos mustelus). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 10.10.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the viral haemorrhagic septicaemia status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to VHS, regardless of the VHS status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 10.10.2, which are intended for any purpose and which comply with Article 5.4.1.:
   a) heat sterilised, hermetically sealed fish products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) pasteurised fish products that have been subjected to a heat treatment at 90°C for at least ten minutes (or to any time/temperature equivalent which has been demonstrated to inactivate VHSV);
   c) mechanically dried, eviscerated fish (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate VHSV);
   d) naturally dried, eviscerated fish (i.e. sun-dried or wind-dried);
   e) fish oil;
   f) fish meal;
   g) fish skin leather.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of species referred to in Article 10.10.2, other than those referred to in point 1 of Article 10.10.3, Competent Authorities should require the conditions prescribed in Articles 10.10.7 to 10.10.12 relevant to the VHS status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of species not covered in Article 10.10.2, but which could reasonably be expected to pose a risk of spread of VHS, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.
Chapter 10.10.- Viral haemorrhagic septicaemia

Article 10.10.4.

Country free from viral haemorrhagic septicaemia

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from VHS if all the areas covered by the shared water bodies are declared countries or zones free from VHS (see Article 10.10.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from VHS if:

1) a country where the species referred to in Article 10.10.2. are present but there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression, as described in the corresponding chapter of the Aquatic Manual, may make a self-declaration of freedom from VHS when basic biosecurity conditions have been continuously met in the country for at least the last ten years;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of VHS;

OR

3) it previously made a self-declaration of freedom from VHS and subsequently lost its disease free status due to the detection of VHS but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of VHS;

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 2 of Article 10.10.5.

Article 10.10.5.

Zone or compartment free from viral haemorrhagic septicaemia

If a zone or compartment extends over more than one country, it can only be declared a VHS free zone or compartment if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from VHS may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) a zone or compartment where the species referred to in Article 10.10.2. are present but there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression, as described in the corresponding chapter of the Aquatic Manual, may be declared free from VHS when basic biosecurity conditions have been continuously met in the zone or compartment for at least the last ten years;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of VHS;
OR
3) it previously made a self-declaration of freedom for a zone from VHS and subsequently lost its disease free status due to the detection of VHS in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of VHS.

Article 10.10.6.

Maintenance of free status

A country, zone or compartment that is declared free from VHS following the provisions of point 1 of Articles 10.10.4. or 10.10.5. (as relevant) may maintain its status as free from VHS provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from VHS following the provisions of point 2 of Articles 10.10.4. or 10.10.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from VHS provided that conditions that are conducive to clinical expression of VHS, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of VHS, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 10.10.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from viral haemorrhagic septicaemia

When importing aquatic animals and aquatic animal products of species referred to in Article 10.10.2. from a country, zone or compartment declared free from VHS, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 10.10.4. or 10.10.5. (as applicable) and 10.10.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from VHS.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 10.10.3.

Article 10.10.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from viral haemorrhagic septicaemia

When importing for aquaculture, aquatic animals of a species referred to in Article 10.10.2. from a country, zone or compartment not declared free from VHS, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive VHSV in accordance with Chapters 4.3., 4.7. and 5.5.
Chapter 10.10.- Viral haemorrhagic septicaemia

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for VHS.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for VHSV in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of VHS (as described in Chapter 2.3.10. of the Aquatic Manual) and test for VHSV in accordance with Chapter 1.4.;
      v) if VHSV is not detected in the F-1 population, it may be defined as free from VHS and may be released from quarantine;
      vi) if VHSV is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 10.10.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from viral haemorrhagic septicaemia

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 10.10.2. from a country, zone or compartment not declared free from VHS, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 10.10.3., or products described in point 1 of Article 10.10.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of VHSV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

Article 10.10.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use from a country, zone or compartment not declared free from viral haemorrhagic septicaemia

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 10.10.2. from a country, zone or compartment not declared free from VHS, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of VHSV.

This article does not apply to commodities referred to in point 1 of Article 10.10.3.
Chapter 10.10.- Viral haemorrhagic septicaemia

Article 10.10.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from viral haemorrhagic septicaemia

1) Competent Authorities should not require any conditions related to VHS, regardless of the VHS status of the exporting country, zone or compartment, when authorising the importation or transit of fish fillets or steaks (chilled or frozen) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 10.10.2. from a country, zone or compartment not declared free from VHS, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

Article 10.10.12.

Importation of disinfected eggs for aquaculture from a country, zone or compartment not declared free from viral haemorrhagic septicaemia

1) When importing disinfected eggs of the species referred to in Article 10.10.2. for aquaculture, from a country, zone or compartment not declared free from VHS, the Competent Authority of the importing country should assess the risk associated with at least:

   a) the VHS virus status of the water to be used during the disinfection of the eggs;
   b) the prevalence of infection with VHS virus in broodstock (ovarian fluid and milt); and
   c) the temperature and pH of the water to be used for disinfection.

2) If the Competent Authority of the importing country concludes that the importation is acceptable, it should apply the following risk mitigation measures including:

   a) the eggs should be disinfected prior to importing, in accordance with recommendations in Chapter 4.4. or those specified by the Competent Authority of the importing country; and
   b) between disinfection and the import, eggs should not come into contact with anything which may affect their health status.

The Competent Authority may wish to consider internal measures, such as renewed disinfection of the eggs upon arrival in the importing country.

3) When importing disinfected eggs of the species referred to in Article 10.10.2. for aquaculture, from a country, zone or compartment not declared free from VHS, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that the procedures described in point 2 of Article 10.10.12. have been fulfilled.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
SECTION 11.
DISEASES OF MOLLUSCS

CHAPTER 11.1.
INFECTION WITH ABALONE HERPESVIRUS

Article 11.1.1.
For the purposes of the Aquatic Code, infection with abalone herpesvirus (AbHV) means infection with the herpesvirus known to cause disease in abalone.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 11.1.2.
Scope

The recommendations in this chapter apply to: Haliotis diversicolor (subspecies aquatilis and supertexta), Haliotis laevegata, Haliotis rubra and hybrids of Haliotis laevegata x Haliotis rubra. These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 11.1.3.
Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with abalone herpesvirus status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with abalone herpesvirus regardless of the infection with abalone herpesvirus status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animal products from the species referred to in Article 11.1.2. which are intended for any purpose and which comply with Article 5.4.1.:
   a) heat sterilised hermetically sealed abalone products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent);
   b) mechanically dried abalone products (i.e. a heat treatment at 100°C for at least 30 minutes or any time/temperature equivalent which has been demonstrated to inactivate AbHV).

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 11.1.2., other than those referred to in point 1 of Article 11.1.3., Competent Authorities should require the conditions prescribed in Articles 11.1.7. to 11.1.11. relevant to the infection with abalone herpesvirus status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 11.1.2. but which could reasonably be expected to pose a risk of spread of infection with abalone herpesvirus, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.
Country free from infection with abalone herpesvirus

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with abalone herpesvirus if all the areas covered by the shared water bodies are declared countries or zones free from infection with abalone herpesvirus (see Article 11.1.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with abalone herpesvirus if:

1) none of the susceptible species referred to in Article 11.1.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 11.1.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with abalone herpesvirus;

OR

4) it previously made a self-declaration of freedom from infection with abalone herpesvirus and subsequently lost its disease free status due to the detection of infection with abalone herpesvirus but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with abalone herpesvirus.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 11.1.5.

Zone or compartment free from infection with abalone herpesvirus

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with abalone herpesvirus if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with abalone herpesvirus may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 11.1.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;
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OR

2) any of the susceptible species referred to in Article 11.1.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with abalone herpesvirus;

OR

4) it previously made a self-declaration of freedom for a zone from infection with abalone herpesvirus and subsequently lost its disease free status due to the detection of infection with abalone herpesvirus in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with abalone herpesvirus.

Article 11.1.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with abalone herpesvirus following the provisions of points 1 or 2 of Articles 11.1.4. or 11.1.5. (as relevant) may maintain its status as free from infection with abalone herpesvirus provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with abalone herpesvirus following the provisions of point 3 of Articles 11.1.4. or 11.1.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with abalone herpesvirus, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with abalone herpesvirus, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 11.1.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with abalone herpesvirus

When importing aquatic animals and aquatic animal products of species referred to in Article 11.1.2. from a country, zone or compartment declared free from infection with abalone herpesvirus, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 11.1.4. or 11.1.5. (as applicable) and 11.1.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with abalone herpesvirus.
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The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 11.1.3.

Article 11.1.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with abalone herpesvirus

When importing for aquaculture, aquatic animals of a species referred to in Article 11.1.2. from a country, zone or compartment not declared free from infection with abalone herpesvirus, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive abalone herpesvirus in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with abalone herpesvirus.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for abalone herpesvirus in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with abalone herpesvirus (as described in Chapter 2.4.1. of the Aquatic Manual) and test for abalone herpesvirus in accordance with Chapter 1.4.;
      v) if abalone herpesvirus is not detected in the F-1 population, it may be defined as free from infection with abalone herpesvirus and may be released from quarantine;
      vi) if abalone herpesvirus is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 11.1.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with abalone herpesvirus

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 11.1.2. from a country, zone or compartment not declared free from infection with abalone herpesvirus, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 11.1.3., or products described in point 1 of Article 11.1.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of AbHV or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
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Article 11.1.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with abalone herpesvirus

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 11.1.2. from a country, zone or compartment not declared free from infection with abalone herpesvirus, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of AbHV.

This article does not apply to commodities referred to in point 1 of Article 11.1.3.

Article 11.1.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with abalone herpesvirus

1) Competent Authorities should not require any conditions related to infection with abalone herpesvirus, regardless of the infection with abalone herpesvirus status of the exporting country, zone or compartment, when authorising the importation or transit of off the shell and eviscerated abalone meat (chilled or frozen) which have been prepared and packaged for retail trade and which comply with Article 5.4.2. Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 11.1.2. from a country, zone or compartment not declared free from infection with abalone herpesvirus, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

CHAPTER 11.2.

INFECTION WITH BONAMIA EXITIOSA

Article 11.2.1.

For the purposes of the Aquatic Code, infection with Bonamia exitiosa means infection with B. exitiosa.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 11.2.2.

Scope

The recommendations in this chapter apply to: Australian mud oyster (Ostrea angasi) and Chilean flat oyster (Ostrea chilensis). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 11.2.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with B. exitiosa status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with B. exitiosa, regardless of the infection with B. exitiosa status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animals and aquatic animal products from the species referred to in Article 11.2.2. which are intended for any purpose and which comply with Article 5.4.1.:
   a) frozen oyster meat; and
   b) frozen half-shell oysters.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 11.2.2., other than those referred to in point 1 of Article 11.2.3., Competent Authorities should require the conditions prescribed in Articles 11.2.7. to 11.2.11. relevant to the infection with B. exitiosa status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 11.2.2. but which could reasonably be expected to pose a risk of spread of infection with B. exitiosa, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 11.2.4.

Country free from infection with B. exitiosa

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with B. exitiosa if all the areas covered by the shared water bodies are declared countries or zones free from infection with B. exitiosa (see Article 11.2.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with B. exitiosa if:

1) none of the susceptible species referred to in Article 11.2.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;
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OR

2) any of the susceptible species referred to in Article 11.2.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with B. exitiosa;

OR

4) it previously made a self-declaration of freedom from infection with B. exitiosa and subsequently lost its disease free status due to the detection of infection with B. exitiosa but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with B. exitiosa.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 11.2.5.

Article 11.2.5.

Zone or compartment free from infection with B. exitiosa

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with B. exitiosa if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with B. exitiosa may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 11.2.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 11.2.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with B. exitiosa;
OR

4) it previously made a self-declaration of freedom for a zone from infection with *B. exitiosa* and subsequently lost its disease free status due to the detection of infection with *B. exitiosa* in the zone but the following conditions have been met:

a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and

b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and

c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and

d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with *B. exitiosa*.

Article 11.2.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with *B. exitiosa* following the provisions of points 1 or 2 of Articles 11.2.4. or 11.2.5. (as relevant) may maintain its status as free from infection with *B. exitiosa* provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with *B. exitiosa* following the provisions of point 3 of Articles 11.2.4. or 11.2.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with *B. exitiosa*, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with *B. exitiosa*, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 11.2.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with *B. exitiosa*

When importing aquatic animals and aquatic animal products of species referred to in Article 11.2.2. from a country, zone or compartment declared free from infection with *B. exitiosa*, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 11.2.4. or 11.2.5. (as applicable) and 11.2.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with *B. exitiosa*.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 11.2.3.
Article 11.2.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with *B. exitiosa*

When importing for *aquaculture*, *aquatic animals* of a species referred to in Article 11.2.2. from a country, zone or compartment not declared free from infection with *B. exitiosa*, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported *aquatic animals*, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported *aquatic animals* in a *quarantine* facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive *B. exitiosa* in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for *aquaculture*, consider applying the following:
   a) In the *exporting country*:
      i) identify potential source populations and evaluate their *aquatic animal* health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of *aquatic animals* with a high health status for infection with *B. exitiosa*.
   b) In the *importing country*:
      i) import the F-0 population into a *quarantine* facility;
      ii) test the F-0 population for *B. exitiosa* in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in *quarantine*;
      iv) culture F-1 population in *quarantine* under conditions that are conducive to the clinical expression of infection with *B. exitiosa* (as described in Chapter 2.4.2. of the *Aquatic Manual*) and test for *B. exitiosa* in accordance with Chapter 1.4.;
      v) if *B. exitiosa* is not detected in the F-1 population, it may be defined as free from infection with *B. exitiosa* and may be released from *quarantine*;
      vi) if *B. exitiosa* is detected in the F-1 population, those animals should not be released from *quarantine* and should be killed and disposed of in a biosecure manner.

Article 11.2.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with *B. exitiosa*

When importing, for processing for human consumption, *aquatic animals* or *aquatic animal products* of species referred to in Article 11.2.2. from a country, zone or compartment not declared free from infection with *B. exitiosa*, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in *quarantine* or containment facilities until processing into one of the products referred to in point 1 of Article 11.2.3., or products described in point 1 of Article 11.2.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of *B. exitiosa* or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 11.2.- Infection with Bonamia exitiosa

Article 11.2.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with B. exitiosa

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 11.2.2. from a country, zone or compartment not declared free from infection with B. exitiosa, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of B. exitiosa.

This article does not apply to commodities referred to in point 1 of Article 11.2.3.

Article 11.2.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with B. exitiosa

1) Competent Authorities should not require any conditions related to infection with B. exitiosa, regardless of the infection with B. exitiosa status of the exporting country, zone or compartment, when authorising the importation or transit of the following commodities which have been prepared and packaged for retail trade and which comply with Article 5.4.2.:

   a) chilled oyster meat; and

   b) chilled half-shell oysters.

Certain assumptions have been made in assessing the safety of the aquatic animal products listed above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 11.2.2. from a country, zone or compartment not declared free from infection with B. exitiosa, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2003; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 11.3.

INFECTION WITH BONAMIA OSTREAE

Article 11.3.1.

For the purposes of the Aquatic Code, infection with Bonamia ostreae means infection with B. ostreae.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 11.3.2.

Scope

The recommendations in this chapter apply to: European flat oyster (Ostrea edulis), Australian mud oyster (Ostrea angasi), Argentinean flat oyster (Ostrea puelchana), Chilean flat oyster (Ostrea chilensis), Asiatic oyster (Ostrea denselammellosa) and Suminoe oyster (Crassostrea ariakensis). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 11.3.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with B. ostreae status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with B. ostreae, regardless of the infection with B. ostreae status of the exporting country, zone or compartment, when authorising the importation or transit of the following aquatic animals and aquatic animal products from the species referred to in Article 11.3.2. which are intended for any purpose and which comply with Article 5.4.1.:
   a) frozen oyster meat;
   b) frozen half-shell oysters.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 11.3.2., other than those referred to in point 1 of Article 11.3.3., Competent Authorities should require the conditions prescribed in Articles 11.3.7. to 11.3.11. relevant to the infection with B. ostreae status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 11.3.2., but which could reasonably be expected to pose a risk of spread of infection with B. ostreae, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 11.3.4.

Country free from infection with B. ostreae

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with B. ostreae if all the areas covered by the shared water bodies are declared countries or zones free from infection with B. ostreae (see Article 11.3.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with B. ostreae if:

1) none of the susceptible species referred to in Article 11.3.2. are present and basic biosecurity conditions have been continuously met for at least the last two years;
Chapter 11.3.- Infection with Bonamia ostreae

OR

2) any of the susceptible species referred to in Article 11.3.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with B. ostreae;

OR

4) it previously made a self-declaration of freedom from infection with B. ostreae and subsequently lost its disease free status due to the detection of infection with B. ostreae but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with B. ostreae.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 11.3.5.

Article 11.3.5.

Zone or compartment free from infection with B. ostreae

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with B. ostreae if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with B. ostreae may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 11.3.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last two years;

OR

2) any of the susceptible species referred to in Article 11.3.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last two years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last two years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with B. ostreae;
Chapter 11.3.- Infection with Bonamia ostreae

OR

4) it previously made a self-declaration of freedom for a zone from infection with B. ostreae and subsequently lost its disease free status due to the detection of infection with B. ostreae in the zone but the following conditions have been met:

a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and

b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and

c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and

d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with B. ostreae.

Article 11.3.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with B. ostreae following the provisions of points 1 or 2 of Articles 11.3.4. or 11.3.5. (as relevant) may maintain its status as free from infection with B. ostreae provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with B. ostreae following the provisions of point 3 of Articles 11.3.4. or 11.3.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with B. ostreae provided that conditions that are conducive to clinical expression of infection with B. ostreae, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with B. ostreae, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 11.3.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with B. ostreae

When importing aquatic animals and aquatic animal products of species referred to in Article 11.3.2. from a country, zone or compartment declared free from infection with B. ostreae, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 11.3.4. or 11.3.5. (as applicable) and 11.3.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with B. ostreae.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 11.3.3.
Chapter 11.3.- Infection with Bonamia ostreae

Article 11.3.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with B. ostreae

When importing for aquaculture, aquatic animals of a species referred to in Article 11.3.2. from a country, zone or compartment not declared free from infection with B. ostreae, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive B. ostreae in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with B. ostreae.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for B. ostreae in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with B. ostreae (as described in Chapter 2.4.3. of the Aquatic Manual) and test for B. ostreae in accordance with Chapter 1.4.;
      v) if B. ostreae is not detected in the F-1 population, it may be defined as free from infection with B. ostreae and may be released from quarantine;
      vi) if B. ostreae is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 11.3.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with B. ostreae

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 11.3.2. from a country, zone or compartment not declared free from infection with B. ostreae, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 11.3.3., or products described in point 1 of Article 11.3.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of B. ostreae or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 11.3.- Infection with Bonamia ostreae

Article 11.3.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with B. ostreae

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 11.3.2. from a country, zone or compartment not declared free from infection with B. ostreae, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of B. ostreae.

This article does not apply to commodities referred to in point 1 of Article 11.3.3.

Article 11.3.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with B. ostreae

1) Competent Authorities should not require any conditions related to infection with B. ostreae, regardless of the infection with B. ostreae status of the exporting country, zone or compartment, when authorising the importation or transit of the following commodities which have been prepared and packaged for retail trade and which comply with Article 5.4.2.:

a) chilled oyster meat;

b) chilled half-shell oysters.

Certain assumptions have been made in assessing the safety of the aquatic animal products listed above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 11.3.2. from a country, zone or compartment not declared free from infection with B. ostreae, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 11.4.

INFECTION WITH MARTEILIA REFRINGENS

Article 11.4.1.

For the purposes of the Aquatic Code, infection with Marteilia refringens means infection with M. refringens.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 11.4.2.

Scope

The recommendations in this chapter apply to: European flat oyster (Ostrea edulis), Australian mud oyster (Ostrea angasi), Argentinean oyster (Ostrea puelchana), Chilean flat oyster (Ostrea chilensis), blue mussel (Mytilus edulis) and Mediterranean mussel (Mytilus galloprovincialis). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 11.4.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with M. refringens status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with M. refringens, regardless of the infection with M. refringens status of the exporting country, zone or compartment, when authorising the importation or transit of heat sterilised hermetically sealed mollusc products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent) from the species referred to in Article 11.4.2. which are intended for any purpose and which comply with Article 5.4.1.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 11.4.2., other than those referred to in point 1 of Article 11.4.3., Competent Authorities should require the conditions prescribed in Articles 11.4.7. to 11.4.11. relevant to the infection with M. refringens status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 11.4.2. but which could reasonably be expected to pose a risk of spread of infection with M. refringens, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 11.4.4.

Country free from infection with M. refringens

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with M. refringens if all the areas covered by the shared water bodies are declared countries or zones free from infection with M. refringens (see Article 11.4.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with M. refringens if:

1) none of the susceptible species referred to in Article 11.4.2. are present and basic biosecurity conditions have been continuously met for at least the last three years;
Chapter 11.4.- Infection with Marteilia refringens

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 11.4.5.

Article 11.4.5.

Zone or compartment free from infection with M. refringens

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with M. refringens if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with M. refringens may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 11.4.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last three years;

OR

2) any of the susceptible species referred to in Article 11.4.2. are present in the zone or compartment and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last three years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last three years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two of the last three years without detection of infection with M. refringens;
Chapter 11.4. - Infection with Marteilia refringens

OR

4) it previously made a self-declaration of freedom for a zone from infection with *M. refringens* and subsequently lost its disease free status due to the detection of infection with *M. refringens* in the zone but the following conditions have been met:

   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and

   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and

   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and

   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two of the last three years without detection of infection with *M. refringens*.

Article 11.4.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with *M. refringens* following the provisions of points 1 or 2 of Articles 11.4.4. or 11.4.5. (as relevant) may maintain its status as free from infection with *M. refringens* provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with *M. refringens* following the provisions of point 3 of Articles 11.4.4. or 11.4.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with *M. refringens*, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with *M. refringens*, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 11.4.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with *M. refringens*

When importing aquatic animals and aquatic animal products of species referred to in Article 11.4.2. from a country, zone or compartment declared free from infection with *M. refringens*, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 11.4.4. or 11.4.5. (as applicable) and 11.4.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with *M. refringens*.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 11.4.3.
Chapter 11.4. - Infection with Marteilia refringens

Article 11.4.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with *M. refringens*

When importing for *aquaculture*, aquatic animals of a species referred to in Article 11.4.2. from a country, zone or compartment not declared free from infection with *M. refringens*, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive *M. refringens* in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for *aquaculture*, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with *M. refringens*.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for *M. refringens* in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with *M. refringens* (as described in Chapter 2.4.4. of the Aquatic Manual) and test for *M. refringens* in accordance with Chapter 1.4.;
      v) if *M. refringens* is not detected in the F-1 population, it may be defined as free from infection with *M. refringens* and may be released from quarantine;
      vi) if *M. refringens* is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 11.4.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with *M. refringens*

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 11.4.2. from a country, zone or compartment not declared free from infection with *M. refringens*, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 11.4.3., or products described in point 1 of Article 11.4.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of *M. refringens* or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.
Chapter 11.4.- Infection with Marteilia refringens

Article 11.4.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with *M. refringens*

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 11.4.2. from a country, zone or compartment not declared free from infection with *M. refringens*, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of *M. refringens*.

This article does not apply to commodities referred to in point 1 of Article 11.4.3.

Article 11.4.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with *M. refringens*

1) Competent Authorities should not require any conditions related to infection with *M. refringens*, regardless of the infection with *M. refringens* status of the exporting country, zone or compartment, when authorising the importation or transit of the following commodities which have been prepared and packaged for retail trade and which comply with Article 5.4.2.:

   a) mollusc meat (chilled or frozen); and
   b) half-shell oysters (chilled or frozen).

Certain assumptions have been made in assessing the safety of the aquatic animal products listed above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 11.4.2. from a country, zone or compartment not declared free from infection with *M. refringens*, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 11.5.

INFECTION WITH PERKINSUS MARINUS

Article 11.5.1.

For the purposes of the Aquatic Code, infection with Perkinsus marinus means infection with P. marinus.

Information on methods for diagnosis is provided in the Aquatic Manual.

Article 11.5.2.

Scope

The recommendations in this chapter apply to: Eastern oyster (Crassostrea virginica), Pacific oyster (Crassostrea gigas), Suminoe oyster (Crassostrea ariakensis), soft shell clam (Mya arenaria), Baltic clam (Macoma balthica) and hard shell clam (Mercenaria mercenaria). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 11.5.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with P. marinus status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with P. marinus, regardless of the infection with P. marinus status of the exporting country, zone or compartment, when authorising the importation or transit of heat sterilised hermetically sealed mollusc products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent) from the species referred to in Article 11.5.2. which are intended for any purpose and which comply with Article 5.4.1.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 11.5.2., other than those referred to in point 1 of Article 11.5.3., Competent Authorities should require the conditions prescribed in Articles 11.5.7. to 11.5.11. relevant to the infection with P. marinus status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 11.5.2. but which could reasonably be expected to pose a risk of spread of infection with P. marinus, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 11.5.4.

Country free from infection with P. marinus

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with P. marinus if all the areas covered by the shared water bodies are declared countries or zones free from infection with P. marinus (see Article 11.5.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with P. marinus if:

1) none of the susceptible species referred to in Article 11.5.2. are present and basic biosecurity conditions have been continuously met for at least the last three years;
Chapter 11.5. - Infection with Perkinsus marinus

2) any of the susceptible species referred to in Article 11.5.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last three years;

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last three years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last three years without detection of infection with P. marinus;

4) it previously made a self-declaration of freedom from infection with P. marinus and subsequently lost its disease free status due to the detection of infection with P. marinus but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last three years without detection of infection with P. marinus.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 11.5.5.

Article 11.5.5.

Zone or compartment free from infection with P. marinus

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with P. marinus if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with P. marinus may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 11.5.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last three years;

2) any of the susceptible species referred to in Article 11.5.2. are present in the zone or compartment and the following conditions have been met:
   a) there has not been any observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last three years;

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last three years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last three years without detection of infection with P. marinus;
Chapter 11.5.- Infection with Perkinsus marinus

OR

4) it previously made a self-declaration of freedom from infection with P. marinus and subsequently lost its disease free status due to the detection of infection with P. marinus but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last three years without detection of infection with P. marinus.

Article 11.5.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with P. marinus following the provisions of points 1 or 2 of Articles 11.5.4. or 11.5.5. (as relevant) may maintain its status as free from infection with P. marinus provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with P. marinus following the provisions of point 3 of Articles 11.5.4. or 11.5.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with P. marinus provided that conditions that are conducive to clinical expression of infection with P. marinus, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with P. marinus, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 11.5.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with P. marinus

When importing aquatic animals and aquatic animal products of species referred to in Article 11.5.2. from a country, zone or compartment declared free from infection with P. marinus, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 11.5.4. or 11.5.5. (as applicable) and 11.5.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with P. marinus.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 11.5.3.
Chapter 11.5.- Infection with Perkinsus marinus

Article 11.5.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with *P. marinus*

When importing for *aquaculture, aquatic animals* of a species referred to in Article 11.5.2. from a country, zone or *compartment* not declared free from infection with *P. marinus*, the *Competent Authority* of the *importing country* should assess the *risk* in accordance with Chapter 2.1. and consider the *risk* mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported *aquatic animals*, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported *aquatic animals* in a *quarantine* facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive *P. marinus* in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for *aquaculture*, consider applying the following:
   a) In the *exporting country*:
      i) identify potential source populations and evaluate their *aquatic animal* health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of *aquatic animals* with a high health status for infection with *P. marinus*.
   b) In the *importing country*:
      i) import the F-0 population into a *quarantine* facility;
      ii) test the F-0 population for ranavirus in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in *quarantine*;
      iv) culture F-1 population in *quarantine* under conditions that are conducive to the clinical expression of infection with *P. marinus* (as described in Chapter 2.4.6. of the *Aquatic Manual*) and test for *P. marinus* in accordance with Chapter 1.4.;
      v) if *P. marinus* is not detected in the F-1 population, it may be defined as free from infection with *P. marinus* and may be released from *quarantine*;
      vi) if *P. marinus* is detected in the F-1 population, those animals should not be released from *quarantine* and should be killed and disposed of in a biosecure manner.

Article 11.5.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with *P. marinus*

When importing, for processing for human consumption, *aquatic animals* or *aquatic animal products* of species referred to in Article 11.5.2. from a country, zone or *compartment* not declared free from infection with *P. marinus*, the *Competent Authority* of the *importing country* should assess the *risk* and, if justified, require that:

1) the consignment is delivered directly to and held in *quarantine* or containment facilities until processing into one of the products referred to in point 1 of Article 11.5.3., or products described in point 1 of Article 11.5.11., or other products authorised by the *Competent Authority*; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of *P. marinus* or is disposed in a manner that prevents contact of waste with *susceptible species*.

For these *commodities* Member Countries may wish to consider introducing internal measures to address the *risks* associated with the *commodity* being used for any purpose other than for human consumption.
Chapter 11.5.- Infection with *Perkinsus marinus*

Article 11.5.10.

**Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with *P. marinus***

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 11.5.2. from a country, zone or compartment not declared free from infection with *P. marinus*, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of *P. marinus*.

This article does not apply to commodities referred to in point 1 of Article 11.5.3.

Article 11.5.11.

**Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with *P. marinus***

1) **Competent Authorities** should not require any conditions related to infection with *P. marinus*, regardless of the infection with *P. marinus* status of the exporting country, zone or compartment, when authorising the importation or transit of the following commodities which have been prepared and packaged for retail trade and which comply with Article 5.4.2.:

   a) mollusc meat (chilled and frozen); and
   b) half-shell oysters (chilled and frozen).

Certain assumptions have been made in assessing the safety of the aquatic animal products listed above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities, Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 11.5.2. from a country, zone or compartment not declared free from infection with *P. marinus*, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

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**NB:** FIRST ADOPTED IN 2000; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 11.6.

INFECTION WITH PERKINSUS OLSENI

Article 11.6.1.

For the purposes of the Aquatic Code, infection with Perkinsus olseni means infection with *P. olseni*. Information on methods for diagnosis are provided in the Aquatic Manual.

Article 11.6.2.

Scope

The recommendations in this chapter apply to: primarily venerid clams (*Austrovenus stutchburyi, Venerupis pullastra, Venerupis aurea, Ruditapes decussatus* and *Ruditapes philippinarum*), abalone (*Haliotis rubra, Haliotis laevigata, Haliotis Cyclobates* and *Haliotis scalaris*) and other species (*Anadara trapezia, Barbatia novaezelandiae, Macomona liliana, Paphies australis* and *Crassostrea ariakensis*). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 11.6.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with *P. olseni* status of the exporting country, zone or compartment

1) *Competent Authorities* should not require any conditions related to infection with *P. olseni*, regardless of the infection with *P. olseni* status of the exporting country, zone or compartment, when authorising the importation or transit of heat sterilised hermetically sealed mollusc products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent) from the species referred to in Article 11.6.2. which are intended for any purpose and which comply with Article 5.4.1.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 11.6.2., other than those referred to in point 1 of Article 11.6.3., *Competent Authorities* should require the conditions prescribed in Articles 11.6.7. to 11.6.11. relevant to the infection with *P. olseni* status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 11.6.2. but which could reasonably be expected to pose a risk of spread of infection with *P. olseni*, the *Competent Authority* should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The *Competent Authority* of the exporting country should be informed of the outcome of this assessment.

Article 11.6.4.

Country free from infection with *P. olseni*

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with *P. olseni* if all the areas covered by the shared water bodies are declared countries or zones free from infection with *P. olseni* free (see Article 11.6.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with *P. olseni* if:

1) a country where the susceptible species referred to in Article 11.6.2. are present but there has been no observed occurrence of the disease for at least the last ten years despite conditions – in all areas where the species are present – that are conducive to its clinical expression, as described in the corresponding chapter of the Aquatic Manual, may make a self-declaration of freedom from infection with *P. olseni* when basic biosecurity conditions have been continuously met in the country for at least the last three years and infection with *P. olseni* is not known to be established in wild populations;
Chapter 11.6. - Infection with Perkinsus olseni

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last three years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last three years without detection of infection with P. olseni;

OR

3) it previously made a self-declaration of freedom from infection with P. olseni and subsequently lost its disease free status due to the detection of infection with P. olseni but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last three years without detection of infection with P. olseni.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 2 of Article 11.6.5.

Article 11.6.5.

Zone or compartment free from infection with P. olseni

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with P. olseni if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with P. olseni may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) in a country of unknown status for infection with P. olseni, a zone or compartment where the susceptible species referred to in Article 11.6.2. are present but there has been no observed occurrence of the disease for at least the last ten years despite conditions – in all areas where the species are present – that are conducive to its clinical expression, as described in the corresponding chapter of the Aquatic Manual, may be declared free from infection with P. olseni when basic biosecurity conditions have been continuously met in the zone or compartment for at least the last three years and infection with P. olseni is not known to be established in wild populations;

OR

2) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last three years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last three years without detection of infection with P. olseni;

OR

3) it previously made a self-declaration of freedom for a zone from infection with P. olseni and subsequently lost its disease free status due to the detection of infection with P. olseni in the zone but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last three years without detection of infection with P. olseni.
Chapter 11.6.- Infection with Perkinsus olseni

Article 11.6.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with *P. olseni* following the provisions of point 1 of Articles 11.6.4. or 11.6.5. (as relevant) may maintain its status as free from infection with *P. olseni* provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with *P. olseni* following the provisions of point 2 of Articles 11.6.4. or 11.6.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with *P. olseni* provided that conditions that are conducive to clinical expression of infection with *P. olseni*, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with *P. olseni*, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 11.6.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with *P. olseni*

When importing aquatic animals and aquatic animal products of species referred to in Article 11.6.2. from a country, zone or compartment declared free from infection with *P. olseni*, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 11.6.4. or 11.6.5. (as applicable) and 11.6.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with *P. olseni*.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 11.6.3.

Article 11.6.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with *P. olseni*

When importing for aquaculture, aquatic animals of a species referred to in Article 11.6.2. from a country, zone or compartment not declared free from infection with *P. olseni*, the Competent Authority of the importing country should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive *P. olseni* in accordance with Chapters 4.3., 4.7. and 5.5.
If the intention is to establish a new stock for aquaculture, consider applying the following:

a) In the exporting country:
   i) identify potential source populations and evaluate their aquatic animal health records;
   ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with P. olseni.

b) In the importing country:
   i) import the F-0 population into a quarantine facility;
   ii) test the F-0 population for P. olseni in accordance with Chapter 1.4. to determine their suitability as broodstock;
   iii) produce a first generation (F-1) population in quarantine;
   iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with P. olseni (as described in Chapter 2.4.7. of the Aquatic Manual) and test for P. olseni in accordance with Chapter 1.4.;
   v) if P. olseni is not detected in the F-1 population, it may be defined as free from infection with P. olseni and may be released from quarantine;
   vi) if P. olseni is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 11.6.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with P. olseni

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 11.6.2. from a country, zone or compartment not declared free from infection with P. olseni, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 11.6.3., or products described in point 1 of Article 11.6.11., or other products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of P. olseni or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

Article 11.6.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with P. olseni

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 11.6.2. from a country, zone or compartment not declared free from infection with P. olseni, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and

2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of P. olseni.

This article does not apply to commodities referred to in point 1 of Article 11.6.3.
Article 11.6.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with *P. olseni*

1) Competent Authorities should not require any conditions related to infection with *P. olseni*, regardless of the infection with *P. olseni* status of the exporting country, zone or compartment, when authorising the importation or transit of the following commodities which have been prepared and packaged for retail trade and which comply with Article 5.4.2.:

   a) mollusc meat (chilled and frozen); and  
   b) half-shell molluscs (chilled and frozen).

Certain assumptions have been made in assessing the safety of the aquatic animal products listed above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 11.6.2. from a country, zone or compartment not declared free from infection with *P. olseni*, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2001; MOST RECENT UPDATE ADOPTED IN 2017.
CHAPTER 11.7.

INFECTION WITH XENOHALIOTIS CALIFORNIENSIS

Article 11.7.1.

For the purposes of the Aquatic Code, infection with Xenohaliotis californiensis means infection with X. californiensis.

Information on methods for diagnosis are provided in the Aquatic Manual.

Article 11.7.2.

Scope

The recommendations in this chapter apply to: black abalone (Haliotis cracherodii), white abalone (Haliotis sorenseni), red abalone (Haliotis rufescens), pink abalone (Haliotis corrugata), green abalone (Haliotis tuberculata and Haliotis fulgens), flat abalone (Haliotis wallalensis) and Japanese abalone (Haliotis discus-hannah). These recommendations also apply to any other susceptible species referred to in the Aquatic Manual when traded internationally.

Article 11.7.3.

Importation or transit of aquatic animals and aquatic animal products for any purpose regardless of the infection with X. californiensis status of the exporting country, zone or compartment

1) Competent Authorities should not require any conditions related to infection with X. californiensis, regardless of the infection with X. californiensis status of the exporting country, zone or compartment, when authorising the importation or transit of heat sterilised hermetically sealed abalone products (i.e. a heat treatment at 121°C for at least 3.6 minutes or any time/temperature equivalent) from the species referred to in Article 11.7.2. which are intended for any purpose and which comply with Article 5.4.1.

2) When authorising the importation or transit of aquatic animals and aquatic animal products of a species referred to in Article 11.7.2., other than those referred to in point 1 of Article 11.7.3., Competent Authorities should require the conditions prescribed in Articles 11.7.7. to 11.7.11. relevant to the infection with X. californiensis status of the exporting country, zone or compartment.

3) When considering the importation or transit of aquatic animals and aquatic animal products of a species not covered in Article 11.7.2. but which could reasonably be expected to pose a risk of spread of infection with X. californiensis, the Competent Authority should conduct a risk analysis in accordance with the recommendations in Chapter 2.1. The Competent Authority of the exporting country should be informed of the outcome of this assessment.

Article 11.7.4.

Country free from infection with X. californiensis

If a country shares a zone with one or more other countries, it can only make a self-declaration of freedom from infection with X. californiensis if all the areas covered by the shared water bodies are declared countries or zones free from infection with X. californiensis (see Article 11.7.5.).

As described in Article 1.4.6., a country may make a self-declaration of freedom from infection with X. californiensis if:

1) none of the susceptible species referred to in Article 11.7.2. are present and basic biosecurity conditions have been continuously met for at least the last three years;
Chapter 11.7. - Infection with Xenohaliotis californiensis

2) any of the susceptible species referred to in Article 11.7.2. are present and the following conditions have been met:
   a) there has been no observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last three years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last three years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with X. californiensis;

OR

4) it previously made a self-declaration of freedom from infection with X. californiensis and subsequently lost its disease free status due to the detection of infection with X. californiensis but the following conditions have been met:
   a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and
   b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and
   c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and
   d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with X. californiensis.

In the meantime, part or all of the non-affected area may be declared a free zone provided that such a part meets the conditions in point 3 of Article 11.7.5.

Article 11.7.5.

Zone or compartment free from infection with X. californiensis

If a zone or compartment extends over more than one country, it can only be declared a zone or compartment free from infection with X. californiensis if all the relevant Competent Authorities confirm that all relevant conditions have been met.

As described in Article 1.4.6., a zone or compartment within the territory of one or more countries not declared free from infection with X. californiensis may be declared free by the Competent Authority(ies) of the country(ies) concerned if:

1) none of the susceptible species referred to in Article 11.7.2. are present in the zone or compartment and basic biosecurity conditions have been continuously met for at least the last three years;

OR

2) any of the susceptible species referred to in Article 11.7.2. are present in the zone or compartment but the following conditions have been met:
   a) there has not been any observed occurrence of the disease for at least the last ten years despite conditions that are conducive to its clinical expression (as described in the corresponding chapter of the Aquatic Manual); and
   b) basic biosecurity conditions have been continuously met for at least the last three years;

OR

3) the disease status prior to targeted surveillance is unknown but the following conditions have been met:
   a) basic biosecurity conditions have been continuously met for at least the last three years; and
   b) targeted surveillance, as described in Chapter 1.4., has been in place, in the zone or compartment, for at least the last two years without detection of infection with X. californiensis;
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OR

4) it previously made a self-declaration of freedom for a zone from infection with X. californiensis and subsequently lost its disease free status due to the detection of infection with X. californiensis in the zone but the following conditions have been met:

a) on detection of the disease, the affected area was declared an infected zone and a protection zone was established; and

b) infected populations have been destroyed or removed from the infected zone by means that minimise the risk of further spread of the disease, and the appropriate disinfection procedures (as described in Chapter 4.3.) have been completed; and

c) previously existing basic biosecurity conditions have been reviewed and modified as necessary and have continuously been in place since eradication of the disease; and

d) targeted surveillance, as described in Chapter 1.4., has been in place for at least the last two years without detection of infection with X. californiensis.

Article 11.7.6.

Maintenance of free status

A country, zone or compartment that is declared free from infection with X. californiensis following the provisions of points 1 or 2 of Articles 11.7.4. or 11.7.5. (as relevant) may maintain its status as free from infection with X. californiensis provided that basic biosecurity conditions are continuously maintained.

A country, zone or compartment that is declared free from infection with X. californiensis following the provisions of point 3 of Articles 11.7.4. or 11.7.5. (as relevant) may discontinue targeted surveillance and maintain its status as free from infection with X. californiensis, as described in the corresponding chapter of the Aquatic Manual, exist and basic biosecurity conditions are continuously maintained.

However, for declared free zones or compartments in infected countries and in all cases where conditions are not conducive to clinical expression of infection with X. californiensis, targeted surveillance needs to be continued at a level determined by the Aquatic Animal Health Service on the basis of the likelihood of infection.

Article 11.7.7.

Importation of aquatic animals and aquatic animal products from a country, zone or compartment declared free from infection with X. californiensis

When importing aquatic animals and aquatic animal products of species referred to in Article 11.7.2. from a country, zone or compartment declared free from infection with X. californiensis, the Competent Authority of the importing country should require that the consignment be accompanied by an international aquatic animal health certificate issued by the Competent Authority of the exporting country or a certifying official approved by the importing country certifying that, on the basis of the procedures described in Articles 11.7.4. or 11.7.5. (as applicable) and 11.7.6., the place of production of the aquatic animals and aquatic animal products is a country, zone or compartment declared free from infection with X. californiensis.

The certificate should be in accordance with the Model Certificate in Chapter 5.11.

This article does not apply to commodities referred to in point 1 of Article 11.7.3.

Article 11.7.8.

Importation of aquatic animals for aquaculture from a country, zone or compartment not declared free from infection with X. californiensis

When importing for aquaculture, aquatic animals of a species referred to in Article 11.7.2. from a country, zone or compartment not declared free from infection with X. californiensis, the Competent Authority of the importing country...
should assess the risk in accordance with Chapter 2.1. and consider the risk mitigation measures in points 1 and 2 below.

1) If the intention is to grow out and harvest the imported aquatic animals, consider applying the following:
   a) the direct delivery to and lifelong holding of the imported aquatic animals in a quarantine facility; and
   b) the treatment of all transport water, equipment, effluent and waste materials to inactive X. californiensis in accordance with Chapters 4.3., 4.7. and 5.5.

OR

2) If the intention is to establish a new stock for aquaculture, consider applying the following:
   a) In the exporting country:
      i) identify potential source populations and evaluate their aquatic animal health records;
      ii) test source populations in accordance with Chapter 1.4. and select a founder population (F-0) of aquatic animals with a high health status for infection with X. californiensis.
   b) In the importing country:
      i) import the F-0 population into a quarantine facility;
      ii) test the F-0 population for X. californiensis in accordance with Chapter 1.4. to determine their suitability as broodstock;
      iii) produce a first generation (F-1) population in quarantine;
      iv) culture F-1 population in quarantine under conditions that are conducive to the clinical expression of infection with X. californiensis (as described in Chapter 2.4.8. of the Aquatic Manual) and test for X. californiensis in accordance with Chapter 1.4.;
      v) if X. californiensis is not detected in the F-1 population, it may be defined as free from infection with X. californiensis and may be released from quarantine;
      vi) if X. californiensis is detected in the F-1 population, those animals should not be released from quarantine and should be killed and disposed of in a biosecure manner.

Article 11.7.9.

Importation of aquatic animals and aquatic animal products for processing for human consumption from a country, zone or compartment not declared free from infection with X. californiensis

When importing, for processing for human consumption, aquatic animals or aquatic animal products of species referred to in Article 11.7.2. from a country, zone or compartment not declared free from infection with X. californiensis, the Competent Authority of the importing country should assess the risk and, if justified, require that:

1) the consignment is delivered directly to and held in quarantine or containment facilities until processing into one of the products referred to in point 1 of Article 11.7.3., or products described in point 1 of Article 11.7.11., or other products authorised by the Competent Authority; and
2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of X. californiensis or is disposed in a manner that prevents contact of waste with susceptible species.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

Article 11.7.10.

Importation of aquatic animals intended for use in animal feed, or for agricultural, industrial or pharmaceutical use, from a country, zone or compartment not declared free from infection with X. californiensis

When importing, for use in animal feed or for agricultural, industrial or pharmaceutical use, aquatic animals of species referred to in Article 11.7.2. from a country, zone or compartment not declared free from infection with X. californiensis, the Competent Authority of the importing country should require that:

1) the consignment is delivered directly to, and held in, quarantine facilities for slaughter and processing into products authorised by the Competent Authority; and
2) water used in transport and all effluent and waste materials from the processing are treated in a manner that ensures inactivation of X. californiensis.
Chapter 11.7.- Infection with Xenohaliotis californiensis

This article does not apply to commodities referred to in point 1 of Article 11.7.3.

Article 11.7.11.

Importation of aquatic animals and aquatic animal products for retail trade for human consumption from a country, zone or compartment not declared free from infection with *X. californiensis*

1) Competent Authorities should not require any conditions related to infection with *X. californiensis*, regardless of the infection with *X. californiensis* status of the exporting country, zone or compartment, when authorising the importation or transit of off the shell, eviscerated abalones (chilled or frozen) which have been prepared and packaged for retail trade and which comply with Article 5.4.2.

Certain assumptions have been made in assessing the safety of the aquatic animal products mentioned above. Member Countries should refer to these assumptions at Article 5.4.2. and consider whether the assumptions apply to their conditions.

For these commodities Member Countries may wish to consider introducing internal measures to address the risks associated with the commodity being used for any purpose other than for human consumption.

2) When importing aquatic animals or aquatic animal products, other than those referred to in point 1 above, of species referred to in Article 11.7.2. from a country, zone or compartment not declared free from infection with *X. californiensis*, the Competent Authority of the importing country should assess the risk and apply appropriate risk mitigation measures.

NB: FIRST ADOPTED IN 2002; MOST RECENT UPDATE ADOPTED IN 2017.
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