## Annex I

### MEETING OF THE OIE

**TERRESTRIAL ANIMAL HEALTH STANDARDS COMMISSION**

**Paris, 2–6 March 2009**

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MEETING OF THE OIE

TERRESTRIAL ANIMAL HEALTH STANDARDS COMMISSION

Paris, 2–6 March 2009

Adopted Agenda

1. Welcome - Director General

2. Update on reports of other commissions and other relevant activities of the OIE - President of the Commission

3. Code revision

A. Examination of Members’ comments

Item 1 Glossary

Item 2 Animal health surveillance (Chapter 1.4.)

Item 3 Horizontal chapters
   a) Import risk analysis (Chapter 2.2.)
   b) Animal health measures applicable before and at departure (Chapter 5.4.)
   c) Border posts and quarantine stations in the importing country (Chapter 5.6.)

Item 4 Design and implementation of animal systems to achieve animal traceability (Chapter 4.2.)

Item 5 Zoning and compartmentalisation
   a) Zoning and compartmentalisation (Chapter 4.3.)
   b) Application of compartmentalisation (Chapter 4.4.)

Item 6 Vector surveillance

Item 7 Semen and embryo (Chapters 4.5., 4.6., 4.7., 4.8., 4.9., 4.10., 4.11.)

Item 8 Somatic cell nuclear transfer in production livestock and horses (Chapter 4.12.)
Annex II (contd)

Item 9  Model certificates
   a) General obligations related to certification (Chapter 5.1.)
   b) Certification procedures (Chapter 5.2.)

Item 10  The role of the Veterinary Services in food safety (Chapter 6.1.)

Item 11  Salmonellosis
   a) Prevention, detection and control of *Salmonella* in poultry (new)
   b) Hygiene and biosecurity procedures in poultry production (Chapter 6.3.)

Item 12  Introduction to the recommendations for controlling antimicrobial resistance (new)

Item 13  Animal welfare
   a) Animal welfare definition (Chapter 7.1.)
   b) Stray dog population control (Member comments on draft chapter)
   c) Report of the *ad hoc* Group on the Welfare of Animals used in Research, Testing and Teaching (laboratory animals)
   d) Report of the *ad hoc* Group on Animal Welfare and Livestock Production Systems
   e) Outcomes of the 2nd global conference on animal welfare

Item 14  Anthrax (Chapter 8.1.)

Item 15  Bluetongue (Chapter 8.3.)

Item 16  Foot and mouth disease (Chapter 8.5.)

Item 17  Leptospirosis (Chapter 8.8.)

Item 18  Paratuberculosis (Chapter 8.10.)

Item 19  Rabies (Chapter 8.11.)

Item 20  Rinderpest (Chapter 8.13.)
Item 21  Bee diseases

a)  Acarapisosis of honey bees (Chapter 9.1.)

b)  American foulbrood of honey bees (Chapter 9.2.)

c)  European foulbrood of honey bees (Chapter 9.3.)

d)  Small hive beetle infestation (Chapter 9.4.)

e)  Tropilaelaps infestation of honey bees (Chapter 9.5.)

f)  Varroosis of honey bees (Chapter 9.6.)

Item 22  Avian influenza (Chapter 10.4.)

Item 23  Newcastle disease (Chapter 10.13)

Item 24  Bovine brucellosis (Chapter 11.3.)

Item 25  Bovine spongiform encephalopathy (Chapter 11.6.)

Item 26  Bovine tuberculosis (Chapter 11.7.) and Bovine tuberculosis of farmed cervidae (new)

Item 27  Contagious bovine pleurpneumonia (Chapter 11.8.)

Item 28  Equine diseases

a)  African horse sickness (Chapter 12.1.)

b)  Equine influenza (Chapter 12.7.)

c)  Equine rhinopneumonitis (Chapter 12.9.)

d)  Equine viral arteritis (Chapter 12.10.)

Item 29  Scrapie (Chapter 14.9.)

Item 30  African swine fever (Chapter 15.1.) and Classical swine fever (Chapter 15.3.)

Item 31  West Nile fever (new)

Item 32  Control of hazards of animal health and public health importance in animal feed
Annex II (contd)

Item 33  Trade in animal products ('Commodities')
   a) Rift Valley fever (Chapter 8.12.)
   b) Bovine cysticercosis (Chapter 11.4.)
   c) Teschovirus encephalomyelitis (Chapter 15.6.)

Item 34  Official disease status questionnaires

Item 35  Infectious bursal disease

Item 36  Swine vesicular disease

B. Other issues

Item 37  Applications for OIE Collaborating Centres (Australia/New Zealand; Uruguay/Chile; Japan)

Item 38  Wildlife disease reporting: Comment on WAHIS

Item 39  OIE Document on the Rights and Obligations of OIE Members in relation to International Trade and Disputes.

Item 40  Future work programme of the Commission

Item 41  Other issues
Glossary

For the purposes of the Terrestrial Code:

**Buffer Protection zone**

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

**Early detection system**

means a system under the control of the Veterinary Services for the timely detection and identification of an incursion or emergence of animal diseases/infections in a country, zone or compartment. Characteristics of the system must include:

- An early detection system should be under the control of the Veterinary Services and should include the following characteristics:
  
  a) representative coverage of target animal populations by field services;
  
  b) ability to undertake effective disease investigation and reporting;
  
  c) access to laboratories capable of diagnosing and differentiating relevant diseases;
  
  d) a training programme for veterinarians, veterinary para-professionals and others involved in handling animals for detecting and reporting unusual disease occurrence;
  
  e) the legal obligation of private veterinarians to report to the Veterinary Authority;
  
  f) a national chain of command.

**Monitoring**

means the intermittent performance and analysis of routine measurements and observations aimed at detecting changes in the environment or health status of a population.

**Official veterinary control of live animals**

means the operations whereby the Veterinary Services, knowing the location of the animals and after taking appropriate actions to identify the identity of their owner or responsible keeper, are able to apply appropriate animal health measures, as required. This does not exclude other responsibilities of the Veterinary Services e.g. food safety.
Annex III (contd)

Outbreak of disease or infection

means the occurrence of one or more cases of a disease or an infection in an epidemiological unit.

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 in the importing country during a specified time period.

Risk assessment

means the evaluation of the likelihood and/or the biological and economic consequences of entry, establishment and spread of a hazard within the territory of an importing country.

Risk communication

is the interactive exchange of information on risk 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.

Sanitary measure

means a measure, such as those described in various Chapters of the Terrestrial Code, destined to protect animal or human health or life within the territory of the OIE Member from risks arising from the entry, establishment and/or spread of a hazard.

Surveillance zone

means a zone established within, and along the border of, a free zone separating the free zone from an infected zone.

The surveillance zone should have an intensified degree of surveillance.

Vaccination

means the successful immunisation of susceptible animals through the administration, according to the manufacturer’s instructions and the Terrestrial Manual, where relevant, of a vaccine comprising antigens appropriate to the disease to be controlled.

Vector

means an insect or any living carrier that transports an infectious agent from an infected individual or its wastes to a susceptible individual or its food or immediate surroundings. The organism may or may not pass through a development cycle within the vector.
Veterinary para-professional

means a person who, for the purposes of the Terrestrial Code, is *authorised* *registered* by the veterinary statutory body to carry out certain designated tasks (dependent upon the category of veterinary para-professional) in a territory, and delegated to them under the responsibility and direction of a veterinarian. The tasks *authorised* for each category of veterinary para-professional should be defined by the veterinary statutory body depending on qualifications and training, and according to need.

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CHAPTER 1.4.

ANIMAL HEALTH SURVEILLANCE

Article 1.4.1.

Introduction and objectives

1. In general, surveillance is aimed at demonstrating the absence of disease or infection, determining the occurrence or distribution of disease or infection, while also detecting as early as possible exotic or emerging diseases. The type of surveillance applied depends on the desired outputs needed to support decision-making. The following recommendations may be applied to all diseases, their agents and all susceptible species (including wildlife) as listed in the Terrestrial Code, and are designed to assist with the development of surveillance methodologies. Except where a specific surveillance method for a certain disease or infection is already described in the Terrestrial Code, the recommendations in this Chapter may be used to further refine the general approaches described for a specific disease or infection. Where detailed disease/infection-specific information is not available, suitable approaches should be based on the recommendations in this Chapter.

2. Animal health surveillance is an essential component necessary tool to detect disease or infection, to monitor disease trends, to facilitate the control of endemic and exotic disease or infection, to support claims for freedom from disease or infection, to provide data to support the for use in risk analysis process, for both animal health and/or public health purposes, and to substantiate the rationale for sanitary measures. Both domestic and wild animals are susceptible to certain disease/infections. However, in the presence of appropriate biosecurity measures, infection/disease/infection in wild animals does not imply that the same infection/disease/infection is necessarily present in domestic animals in the same country or zone. Surveillance data underpin the quality of disease status reports and should satisfy information requirements for accurate risk analysis both for international trade as well as and for national decision-making. Wildlife may be included because these they can serve both as reservoirs and as sensitive indicators of important human and domestic animal, and wildlife diseases. Wildlife disease/infection surveillance presents specific challenges that may differ importantly significantly from disease surveillance in livestock.

3. Essential prerequisites to enable an OIE Member to provide information for the evaluation of its animal health status are:

   a) that the particular Member complies with the provisions of Chapter 3.1. of the Terrestrial Code on the quality and evaluation of the Veterinary 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. of the Terrestrial Code.

4. The objectives of this Chapter are to:

   a) provide guidance to the type of outputs that a surveillance system should generate;

   b) provide recommendations to assess the quality of disease/infection surveillance systems.
Annex IV (contd)

Article 1.4.2.

Definitions

The following definitions apply for the purposes of this Chapter:

**Bias:** A tendency of an estimate to deviate in one direction from a true value.

**Case definition:** A set of criteria used to classify an animal or epidemiological unit as a case.

**Confidence:** In the context of demonstrating freedom from infection, confidence is the probability that the type of surveillance applied would detect the presence of infection if the population were infected. The confidence depends on, among other parameters, the assumed level of infection in an infected population. The term refers to confidence in the ability of the surveillance applied to detect disease, and is equivalent to the sensitivity of the surveillance system.

**Early detection system:** A system for the timely detection and identification of an incursion or emergence of disease in a country, zone or compartment. An early detection system should be under the control of the Veterinary Service and should include the following characteristics:

a) representative coverage of target animal populations by field services;

b) ability to undertake effective disease investigation and reporting;

c) access to laboratories capable of diagnosing and differentiating relevant diseases;

d) a training programme for veterinarians, veterinary para-professionals and others involved in handling animals for detecting and reporting unusual animal health incidents;

e) the legal obligation of private veterinarians in relation to the Veterinary Authority;

f) timely reporting system of the event to the Veterinary Services;

g) a national chain of command.

**Outbreak definition:** An outbreak definition is a set of criteria used to classify the occurrence of one or more cases in a group of animals or units as an outbreak.

**Probability sampling:** A sampling strategy in which every unit has a known non-zero probability of inclusion in the sample.

**Sample:** The group of elements (sampling units) drawn from a population, on which tests are performed or parameters measured to provide surveillance information.

**Sampling units:** The unit that is sampled, either in a random survey or in non-random surveillance. This may be an individual animal or a group of animals (e.g. an epidemiological unit). Together, they comprise the sampling frame.

**Sensitivity:** The proportion of truly positive units that are correctly identified as positive by a test.

**Specificity:** The proportion of truly negative units that are correctly identified as negative by a test.
Study population: The population from which surveillance data are derived. This may be the same as the target population or a subset of it.

Surveillance: The systematic ongoing collection, collation, and analysis of data, and the timely dissemination of information to those who need to know so that action can be taken.

Surveillance system: A method of surveillance that may involve one or more component activities that generates information on the health, disease or zoonosis status of animal populations.

Survey: An investigation in which information is systematically collected, usually carried out on a sample of a defined population group, within a defined time period.

Target population: The population about which conclusions are to be inferred.

Test: A procedure used to classify a unit as either positive, negative or suspect with respect to an infection or disease or infection.

Test system: A combination of multiple tests and rules of interpretation which are used for the same purpose as a test.

Wildlife: Mammals and birds which are not permanently captive or owned free-range. This definition includes the categories of "wild animal" (wild animal genotype living outside of controlling human influence) and "feral animal" (domestic animal genotype living outside of controlling human influence).

Article 1.4.3.

Principles of surveillance

1. Types of surveillance
   
a) Surveillance may be based on many different data sources and can be classified in a number of ways, including:
      
      i) the means by which data are collected (active versus passive surveillance);
      
      ii) the disease focus (pathogen-specific versus general surveillance); and
      
      iii) the way in which units for observation are selected (structured surveys versus non-random data sources).
   
b) In this Chapter, surveillance activities are classified as being based on:
      
      EITHER
      
      i) structured population-based surveys, such as:
         
         - systematic sampling at slaughter;
         
         - random surveys;
         
         - surveys for infection in clinically normal animals, including wildlife;
Annex IV (contd)

OR

ii) structured non-random surveillance activities, such as:
   - disease reporting or notifications;
   - control programmes/health schemes;
   - targeted testing/screening;
   - ante-mortem and post-mortem inspections;
   - laboratory investigation records;
   - biological specimen banks;
   - sentinel units;
   - field observations;
   - farm production records;
   - wildlife disease data.

c) In addition, all available surveillance data should be supported by related information, such as:
   i) data on the epidemiology of the disease/infection, including environmental, host population distribution, and climatic information;
   ii) data on animal movements, and including transhumance, as well as natural wildlife migrations;
   iii) trading patterns for animals and animal products;
   iv) national animal health regulations, including information on compliance with them and their effectiveness;
   v) history of imports of potentially infected material; and
   vi) biosecurity measures in place.

vii) the risk and consequence of disease/infection introduction.

d) 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 structured 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.

2. Critical elements

In assessing the quality of a surveillance system, the following critical elements need to be addressed over and above quality of Veterinary Services (Chapter 3.1.).
a) Populations

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

Definitions of appropriate populations should be based on the specific recommendations of the disease Chapters of the Terrestrial Code.

b) Time frame (or temporal value of surveillance data)

Surveillance should be carried out at a frequency that reflects the biology of the infection and the risks of its introduction.

c) Epidemiological unit

The relevant epidemiological unit for the surveillance system should be defined and documented to ensure that it is representative of the population. Therefore, it should be chosen taking into account factors such as carriers, reservoirs, vectors, immune status, genetic resistance and age, sex, and other host criteria.

d) Clustering

Infection in a country, zone or compartment usually clusters rather than being uniformly or randomly distributed through a population. Clustering may occur at a number of different levels (e.g. a cluster of infected animals within a herd, a cluster of pens in a building, or a cluster of farms in a compartment). Clustering should be taken into account in the design of surveillance activities and the statistical analysis of surveillance data, at least at what is judged to be the most significant level of clustering for the particular animal population and infection.

d) Case and outbreak definitions

A clear and unambiguous case and outbreak definition should be developed and documented for each pathogen disease under surveillance, using, where they exist, the standards in the Terrestrial Code. For wildlife disease / infection surveillance it is essential to correctly identify and report host animal taxonomy (including genus and species).

e) 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 host species, pathogens, varying production and surveillance systems, and types and amounts of data and information available.

The methodology used should be based on the best available information available that is in accord with current scientific thinking. The methodology should also 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.

f) Testing

Surveillance involves the detection of disease or infection by the use of appropriate case definitions based on the results of one or more tests for evidence of infection or immune 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.

The values of sensitivity and specificity for the tests used should be specified for each species in which they may be used, and the method used to determine or estimate these values should be documented. Alternatively, where values for sensitivity and/or specificity for a particular test are specified in the Terrestrial Manual, these values may be used as a guide. For each host species to which a diagnostic test is applied, whenever possible, the tests should be shown to have acceptable sensitivity and specificity for that particular host species.

Samples from a number of 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.

g) 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.

h) 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.

i) 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:

- the distribution of, and communication between, those involved in generating and transferring data from the field to a centralised location; this requires effective collaboration among all stakeholders, such as government ministries, non-governmental organisations, and others, particularly for data involving wildlife.
Annex IV (contd)

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

- maintenance of disaggregated data rather than the compilation summary data;

- minimisation of transcription errors during data processing and communication.

Article 1.4.4.

Structured population-based surveys

In addition to the principles for surveillance discussed above, 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. A sample may be selected in either of the two following ways:

   a) non-probability based sampling methods, such as:

      i) convenience;

      ii) expert choice;

      iii) quota;

   b) probability based sampling methods, such as:

      i) simple random selection;

      ii) cluster sampling;

      iii) stratified sampling;

      iv) systematic sampling.

   Non-probability based sampling methods will not be discussed further.

   Periodic or repeated surveys conducted in order to document disease freedom should be done using probability based sampling methods so that data from the study population can be extrapolated to the target population in a statistically valid manner.

   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 infection and the resources available.
Data on wild animal population size often do not exist and should be determined before a survey can be designed. The expertise of wildlife biologists may be sought in the gathering and interpretation of such population data. Historical population data should be updated since these may not reflect current populations.

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 of interest with respect to the objective of the study such as the presence or absence of infection. 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 an infection in a population of unknown disease status, targeted sampling methods that optimise the detection of infection can be used. In such cases, care should be taken regarding the inferences made from the results.

Specimens from wildlife for surveillance may be available from sources such as hunters and trappers, road-kills, wild animal meat markets, sanitary inspection of hunted animals, morbidity-mortality observations by the general public, wildlife rehabilitation centres, wildlife biologists and wildlife agency field personnel, farmers, and other landholders, naturalists and conservationists. Wildlife data such as census data, trends over time, and reproductive success can be used in a manner similar to farm production records for epidemiological purposes.

4. Sampling methods

When selecting epidemiological units from within a population, probability sampling (e.g. simple random selection) should be used. When this is not possible, sampling should provide the best practical chance of generating a sample that is representative of 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. infection) or to estimate a parameter (e.g. the prevalence of infection). The method used to calculate sample size for surveys depends on the purpose of the survey, the expected prevalence, the level of confidence desired of the survey results and the performance of the tests used.

Article 1.4.5.

Structured non-random surveillance

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

1. Common non-random surveillance sources

A wide variety of non-random surveillance 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 infection. 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). Surveillance systems routinely use structured non-random data, either alone or in combination with surveys.
a) Disease reporting or notification systems

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. Effective laboratory support is an important component of any reporting system. 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 minimized (to hours in the case of introduction of a foreign animal disease).

Whenever the responsibility for disease notification falls outside the scope of the Veterinary Authority, for example in some countries for diseases in wildlife, effective communication and data sharing should be established with the relevant authorities to ensure comprehensive and timely disease reporting.

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 structured surveillance.

c) Targeted testing / screening

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

d) Ante-mortem and post-mortem inspections

Inspections of animals at abattoirs slaughterhouses may provide valuable surveillance data. The sensitivity and specificity of the particular slaughterhouse inspection system for detecting the presence of infectious agents of surveillance interest specified diseases under the particular inspection arrangements applying system in a country place should be pre-determined by the Competent Authority if the data is to be fully utilised. The accuracy of the inspection system will be influenced by:

i) the level of training and experience and number of the inspection staff doing the inspections, and the ratio of staff of different levels of training;

ii) the involvement of the Competent Authorities in the supervision of ante-mortem and post-mortem inspections;

iii) the quality of construction of the abattoir slaughterhouse, speed of the slaughter chain, lighting quality, etc.; and

iv) staff morale and motivation for accurate and efficient performance.

Abattoir Slaughterhouse inspections are likely to provide good coverage only for particular age groups and geographical areas. Abattoir Slaughterhouse surveillance data are subject to obvious biases in relation to target and study populations (e.g. only animals of a particular class and age may are likely to be slaughtered for human consumption in significant numbers). Such biases need to be recognised when analysing surveillance data.
Annex IV (contd)

Both for traceback in the event of detection of disease and for analysis of spatial and herd-level coverage, there should be, if possible, an effective identification system that relates each animal in the abattoir-slaughterhouse 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 standardized methods for interpretation and data recording. As with abattoir inspections, there needs to be a mechanism to relate specimens to the farm of origin.

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 infection, 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/immune status in a specified geographical location to detect the occurrence of disease (usually serologically). 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 infection (related to vector habitats and host population distribution), cost and other practical constraints. Sentinel units may provide evidence of freedom from infection, or provide data on prevalence and incidence as well as the distribution of disease.

h) Field observations

Clinical observations of animals in the field are an important source of surveillance data. The sensitivity and 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 infection at the herd or flock level. In general, the sensitivity of this approach may be quite high (depending on the disease), but the specificity is often quite low.
Wildlife data

Specimens from wild animals for disease/infection surveillance may be available from sources such as hunters and trappers, road-kills, wild animal meat markets, sanitary inspection of hunted animals—mortality and mortality observations by the general public, wildlife rehabilitation centres, wildlife biologists and wildlife agency field personnel, farmers and other landholders, naturalists and conservationists. Wildlife data such as census data, trends over time, and reproductive success can be used in a manner similar to farm production records for epidemiological purposes.

2. Critical elements for structured non-random surveillance

There are a number of critical factors which should be taken into account when using structured 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, however, can be a cost-efficient method of early detection, and may increase the level of confidence or be able to detect a lower level of prevalence with the same level of confidence compared to structured random sampling surveys.

3. Analytical methodologies

Different methodologies may be used for the analysis of non-random surveillance data. Different scientifically valid methodologies may be used for the analysis of non-random surveillance data. Where no 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 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 may provide cumulative evidence of animal health status. Such evidence gathered over time may be combined to provide an overall level of confidence. For instance, repeated annual surveys may be analysed to provide a cumulative 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 just one year.

Analysis of surveillance information gathered intermittently or continuously over time should, where possible, incorporate the time of collection of the information to take the decreased value of older information into account. 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.

Surveillance to demonstrate freedom from disease/infection

1. Requirements to declare a country, zone or compartment free from disease/infection without pathogen specific surveillance

This Article provides general principles for declaring a country, zone or compartment free from disease/infection in relation to the time of last occurrence and in particular for the recognition of historical freedom.
Annex IV (contd)

The provisions of this Article are based on the principles described in Article 1.4.3. of this Chapter and the following premises:

- in the absence of disease and vaccination, the animal population would become susceptible over a period of time;
- the disease agents to which these provisions apply are likely to produce identifiable clinical signs in susceptible animals;
- competent and effective Veterinary Services will be able to investigate, diagnose and report disease, if present;
- disease/infection can affect both wild and domestic animals;
- the absence of disease/infection over a long period of time in a susceptible population can be substantiated by effective disease investigation and reporting by a Member.

a) Historically free

Unless otherwise specified in the relevant disease Chapter, a country, zone or compartment may be recognised free from infection without formally applying a pathogen-specific surveillance programme when:

i) there has never been occurrence of disease, or

ii) eradication has been achieved or the disease/infection has ceased to occur for at least 25 years, provided that for at least the past 10 years:

iii) it has been a notifiable disease;

iv) an early detection system has been in place for all relevant species;

v) measures to prevent disease/infection introduction have been in place; no vaccination against the disease has been carried out unless otherwise provided in the Terrestrial Code;

vi) infection is not known to be established in wildlife 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 infection in wildlife. However, specific surveillance in wildlife is not necessary.)

b) Last occurrence within the previous 25 years

Countries, zones or compartments that have achieved eradication (or in which the disease/infection has ceased to occur) within the previous 25 years, should follow the pathogen-specific surveillance requirements in the Terrestrial Code if they exist. In the absence of specific requirements for surveillance in the Terrestrial Code, countries should follow the general recommendations on surveillance to demonstrate animal health status outlined in this Chapter provided that for at least the past 10 years:

i) it has been a notifiable disease;

ii) an early detection system has been in place;

iii) measures to prevent disease/infection introduction have been in place;
iv) no vaccination against the disease has been carried out unless otherwise provided in the Terrestrial Code;

v) infection is not known to be established in wildlife within the country or zone intended to be declared free. (A country or zone cannot apply for freedom if there is any evidence of infection in wildlife. However, specific surveillance in wildlife is not necessary.)

2. Recommendations for the discontinuation of pathogen-specific screening after recognition of freedom from infection

A country, zone or compartment that has been recognised as free from infection following the provisions of the Terrestrial Code may discontinue pathogen-specific screening while maintaining the infection-free status provided that:

a) it is a notifiable disease;

b) an early detection system is in place;

c) measures to prevent disease/infection introduction are in place;

d) vaccination against the disease is not applied;

e) infection is known not to be established in wildlife. (Specific surveillance in wildlife has demonstrated the absence of infection. It can be difficult to collect sufficient epidemiological data to prove absence of disease/infection in wild animal populations. Therefore, in such circumstances, a range of supporting evidence should be used to make this assessment.)

3. Self declaration of disease/infection

Members may make a self declaration that a country, zone or compartment is free from a listed disease, based on the implementation of the provisions of the Terrestrial Code and the Terrestrial Manual - see relevant provisions in Chapter 1.5. The Veterinary Authority may wish to transmit this information to the OIE Central Bureau, which may publish the information.

4. International recognition of disease/infection free status

For diseases for which procedures exist whereby the OIE can officially recognise the existence of a disease/infection free country, zone or compartment, a Member wishing to apply for recognition of this status shall, via its Permanent Delegate, send to the OIE all the relevant documentation relating to the country, zone or compartment concerned. Such documentation should be presented according to the recommendations prescribed by the OIE for the appropriate animal diseases.

45. Demonstration of freedom from infection

A surveillance system to demonstrate freedom from infection should meet the following requirements in addition to the general requirements for surveillance outlined in Article 1.4.3. of this Chapter.

Freedom from infection implies the absence of the pathogenic agent in the country, zone or compartment. Scientific methods cannot provide absolute certainty of the absence of infection.
Demonstrating freedom from infection involves providing sufficient evidence to demonstrate (to a level of confidence acceptable to Members) that infection 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 infection (unless every member of the population is examined simultaneously with a perfect test with both sensitivity and specificity equal to 100%). Instead, the aim is to provide adequate evidence (to an acceptable level of confidence), that infection, if present, is present in less than a specified proportion of the population.

However, finding evidence of infection at any level in the target population automatically invalidates any freedom from infection claim unless otherwise stated in the relevant disease Chapter. The implications of disease/infection in wildlife for the status of domestic animals in the same country or zone should be assessed in each situation, as indicated in the relevant Chapter on each disease in the Terrestrial Code.

Evidence from targeted, random or non-random data sources, as stated before, may increase the level of confidence or be able to detect a lower level of prevalence with the same level of confidence compared to structured surveys.

**Article 1.4.7.**

**Surveillance for distribution and occurrence of infection**

Surveillance to determine distribution and occurrence of infection or of other relevant health related events is widely used to assess progress in the control or eradication of selected diseases and pathogens and as an aid to decision making. It has, however, relevance for the international movement of animals and products when movement occurs among infected countries.

In contrast to surveillance to demonstrate freedom from infection, surveillance used to assess progress in control or eradication of selected diseases and pathogens is usually designed to collect data about a number of variables of animal health relevance, for example:

1. prevalence or incidence of infection;
2. morbidity and mortality rates;
3. frequency of disease/infection risk factors and their quantification;
4. frequency distribution of herd sizes or the sizes of other epidemiological units;
5. frequency distribution of antibody titres;
6. proportion of immunised animals after a vaccination campaign;
7. frequency distribution of the number of days elapsing between suspicion of infection and laboratory confirmation of the diagnosis and/or to the adoption of control measures;
8. farm production records, etc.
9. role of wildlife in maintenance or transmission of the infection.

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CHAPTER 2.1.

GENERAL CONSIDERATIONS

Article 2.1.1

Introduction

The importation of animals and 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 animals, animal products, animal genetic material, feedstuffs, biological products and pathological material. 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 alludes to the role of the OIE with respect to the Agreement on the Application of Sanitary and Phytosanitary Measures (the so-called SPS Agreement) of the World Trade Organization (WTO), provides definitions and describes the OIE informal procedure for dispute mediation.

Chapter 2.2. provides recommendations and principles for conducting transparent, objective and defensible risk analysis for international trade. The components of risk analysis described in that Chapter 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 which estimates the risks associated with a hazard. Risk assessments may be qualitative or quantitative. For many diseases, particularly for those diseases listed in this Terrestrial 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.
Annex V (contd)

The process of import risk analysis usually needs to take into consideration the results of an evaluation of Veterinary Services, zoning, compartmentalisation and surveillance systems in place for monitoring of animal health in the exporting country. These are described in separate Chapters in the Terrestrial Code.

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CHAPTER 2.2.

IMPORT RISK ANALYSIS

Article 2.2.1.

Introduction

The importation of animals and 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 animals, animal products, animal genetic material, feedstuffs, biological products and pathological material. 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.

Fig 1. The four components of risk analysis

Chapter 2.2. provides recommendations and principles for conducting transparent, objective and defensible risk analyses for international trade. The components of risk analysis described in that Chapter are hazard identification, risk assessment, risk management and risk communication (Figure 1):

The risk assessment is the component of the analysis which estimates the risks associated with a hazard. Risk assessments may be qualitative or quantitative. For many diseases, particularly for those diseases listed in this Terrestrial 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 modeling 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.
Annex V (contd)

The process of import risk analysis usually needs to take into consideration the results of an evaluation of Veterinary Services, zoning, compartmentalisation and surveillance systems in place for monitoring of animal health in the exporting country. These are described in separate Chapters in the Terrestrial Code.

An import risk analysis begins with a description of the commodity proposed for import and the likely annual quantity of trade. It must be recognised that whilst an accurate estimate of the anticipated quantity of trade is desirable to incorporate into the risk estimate, it may not be readily available, particularly where such trade is new.

Hazard identification is an essential step which must be conducted before the risk assessment.

The risk assessment process consists of four interrelated steps. These steps clarify the stages of the risk assessment, describing them in terms of the events necessary for the identified potential risk(s) to occur, and facilitate understanding and evaluation of the outputs. The product is the risk assessment report which is used in risk communication and risk management.

The relationships between risk assessment and risk management processes are outlined in Figure 1.

**Fig 1. The relationship between risk assessment and risk management processes**

**Article 2.2.2.**

**Hazard identification**

The hazard identification involves identifying the pathogenic agents which could potentially produce adverse consequences associated with the importation of a commodity.

The potential 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 potential hazard is already present in the importing country, and whether it is a notifiable 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 potential hazards or not. The risk assessment may be concluded if hazard identification fails to identify potential hazards associated with the importation.
The evaluation of the Veterinary Services, surveillance and control programmes and zoning and compartmentalisation systems are important inputs for assessing the likelihood of hazards being present in the animal population of the exporting country. An importing country may decide to permit the importation using the appropriate sanitary standards recommended in the Terrestrial Code, thus eliminating the need for a risk assessment.

Article 2.2.3.

Principles of risk assessment

1. Risk assessment should be flexible to deal with the complexity of real life situations. No single method is applicable in all cases. Risk assessment must be able to accommodate the variety of 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. Although quantitative assessment is recognised as being able to provide deeper insights into a particular problem, qualitative methods may be more relevant when available data are limited.

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 2.2.4.

Risk assessment steps

1. Release assessment

Release assessment consists of describing the biological pathway(s) necessary for an importation activity to ‘release’ (that is, introduce) pathogenic agents into a particular environment, and estimating the probability of that complete process occurring, either qualitatively (in words) or quantitatively (as a numerical estimate). The release assessment describes the probability of the ‘release’ of each of the potential hazards (the pathogenic agents) 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 release assessment are:

a) Biological factors
   - species, age and breed of animals
   - agent predilection sites
   - vaccination, testing, treatment and quarantine.
b) Country factors
   - incidence/prevalence
   - evaluation of Veterinary Services, surveillance and control programmes and zoning and compartmentalisation systems of the exporting country.

c) Commodity factors
   - quantity of commodity to be imported
   - ease of contamination
   - effect of processing
   - effect of storage and transport.

If the release 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) released from a given risk source, and estimating the probability of the 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 (e.g. ingestion, inhalation, or insect bite), 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.

b) Country factors
   - presence of potential vectors
   - human and animal demographics
   - customs and cultural practices
   - geographical and environmental characteristics.

c) Commodity factors
   - quantity of commodity to be imported
   - intended use of the imported animals or products
   - 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 must 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**
   - animal infection, disease and production losses
   - public health consequences.

b) **Indirect consequences**
   - surveillance and control costs
   - compensation costs
   - potential trade losses
   - adverse consequences to the environment.

4. **Risk estimation**

Risk estimation consists of integrating the results from the release 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:

- estimated numbers of herds, flocks, animals 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.
Principles of risk management

1. Risk management is the process of deciding upon and implementing measures to achieve the Member’s appropriate level of protection, whilst at the same time ensuring that negative effects on trade are minimized. The objective is to manage risk appropriately to ensure that a country’s desire to minimize 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 in the standards.

Risk management components

1. Risk evaluation - the process of comparing the risk estimated in the risk assessment with the Member’s appropriate level of protection.

2. Option evaluation - the process of identifying, evaluating the efficacy and feasibility of, and selecting measures in order to reduce the risk associated with an importation in order to bring i into line with the Members appropriate level of protection. The efficacy is the degree to which an option reduces the likelihood and/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 the 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 and foreign industry groups, domestic livestock producers and consumer 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 is a component of risk communication in order to obtain scientific critique and to ensure that the data, information, methods and assumptions are the best available.

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CHAPTER 5.4.

ANIMAL HEALTH MEASURES APPLICABLE BEFORE AND AT DEPARTURE

Article 5.4.1.

Animals for breeding, rearing or slaughter

1. Countries should only authorise the exportation from their territory of animals for breeding or rearing or animals for slaughter which are correctly identified and which meet the requirements of the importing country.

2. Biological tests and/or vaccinations required by the importing country should be carried out in accordance with the recommendations in the Terrestrial Code and Terrestrial Manual, as well as disinfection and disinestation procedures.

3. Observation of the animals before leaving the country may be carried out either in the establishment where they were reared, or in a quarantine station. When they have been found to be clinically healthy and free from diseases listed by the OIE of the health status agreed by the importing and exporting countries by an Official Veterinarian or by an official of the Competent Authority, during the period of observation, the animals should be transported to the place of shipment in specially constructed vehicles, previously cleansed and if required disinfected if required. This must be done without delay and without the animals coming into contact with other susceptible animals, unless these animals have animal health guarantees similar to those of the transported animals. An international veterinary certificate should attest that the animals have been found to be clinically healthy and of the health status agreed by the importing and exporting countries.

4. The transportation of the animals for breeding or rearing or animals for slaughter from the establishment of origin to the point of departure from the exporting country shall be carried out in conformity with the conditions agreed between the importing country and exporting country.

Article 5.4.2.

Semen, embryo/ova, hatching eggs

Countries should only undertake the export from its territory of:

a) semen,

b) embryos/ova,

c) hatching eggs,

from artificial insemination centres, collection centres or farms which meet the requirements of the importing country.

Article 5.4.3.

Notification

Countries exporting animals, semen, embryos/ova or hatching eggs should inform the country of destination and where necessary the transit countries if, after exportation, a disease listed by the OIE occurs within the incubation period of that particular disease in the establishment of origin, or in an animal which was in an establishment point at which the shipment is assembled of where animals for breeding or rearing or animals for slaughter from different establishments or markets are collected together, or in a market, at the same time as the exported animals.
Article 5.4.4.

Certificate

Before the departure of animals, semen, embryos/ova, hatching eggs and brood-combs of bees, an Official Veterinarian should, within the 24 hours prior to shipment, provide an international veterinary certificate conforming with the models approved by the OIE (as shown in Chapters 5.10. to 5.12. of the Terrestrial Code) and worded in the languages agreed upon between the exporting country and the importing country, and, where necessary, with the transit countries.

Article 5.4.5.

Live animals

1. Before the departure of an animal or a consignment of animals on an international journey, the Veterinary Authority of the port, airport or district in which the border post is situated may, if it is considered necessary, carry out a clinical examination of the animal or consignment. The time and place of the examination shall be arranged taking into account customs and other formalities and in such a way as not to impede or delay departure.

2. The Veterinary Authority referred to in point 1 above shall take necessary measures to:

   a) prevent the shipment of animals affected or suspected of being affected with any disease listed by the OIE or with any other infectious disease as agreed by the importing and exporting countries;

   b) avoid entry into the vehicle of possible vectors or causal agents of infection.

Article 5.4.6.

Products of animal origin

1. Countries should only authorise the export from their territory of meat and products of animal origin intended for human consumption, which are fit for human consumption. They must be accompanied by an international veterinary certificate conforming with the models approved by the OIE (as shown in Chapters 5.10. to 5.12. of the Terrestrial Code). These must be worded in the languages agreed upon between the exporting country and the importing country, and, where necessary, with the transit countries.

2. Products of animal origin intended for use in animal feeding, or for pharmaceutical or surgical or agricultural or industrial use, should be accompanied by an international veterinary certificate conforming with the models approved by the OIE (as shown in Chapters 5.10. to 5.12. of the Terrestrial Code).
CHAPTER 5.6.

BORDER POSTS AND QUARANTINE STATIONS IN THE IMPORTING COUNTRY

Article 5.6.1.

1. Countries and their Veterinary Authorities shall should, wherever possible, take the necessary action to ensure that the border posts and quarantine stations in their territory shall should be provided with an adequate organisation and sufficient equipment for the application of the measures recommended in the Terrestrial Code.

2. Each border post and quarantine station shall should be provided with facilities for the feeding and watering of animals.

Article 5.6.2.

When justified by the amount of international trade and by the epidemiological situation, border posts and quarantine stations shall should be provided with a Veterinary Service comprising personnel, equipment and premises as the case may be and, in particular, means for:

a) making clinical examinations and obtaining specimens of material for diagnostic purposes from live animals or carcasses of animals affected or suspected of being affected by an epizootic disease, and obtaining specimens of animal products suspected of contamination;

b) detecting and isolating animals affected by or suspected of being affected by an epizootic disease;

c) carrying out disinfection and possibly disinfestation of vehicles used to transport animals and animal products.

In addition to this, each port and international airport should ideally be provided with equipment for the sterilisation or incineration of swill or any other material dangerous to animal health.

Article 5.6.3.

When required for the transit of commodities in international trade, airports shall should provide be provided, as soon as possible, with areas of direct transit. These must, however, comply with the conditions required by Veterinary Authorities, especially to prevent contact between animals of different health status and the risk of introducing diseases transmitted by insects.

Article 5.6.4.

Each Veterinary Authority, when requested, shall should make available for the Central Bureau and any interested country on request:

a) a list of border posts, quarantine stations, approved abattoirs and storage depots in its territory which are approved for international trade;
Annex V (contd)

b) the period of time required for notice to be given for the application of the arrangements contained in point 2 of Articles 5.7.1. to 5.7.4.;

c) a list of airports in its territory which are provided with an area of direct transit, approved by the relevant Veterinary Authority and placed under its immediate control, where animals stay for a short time pending further transport to their final destination.

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CHAPTER 4.2.

DESIGN AND IMPLEMENTATION OF IDENTIFICATION SYSTEMS TO ACHIEVE ANIMAL TRACEABILITY

Article 4.2.1.

Introduction and objectives

These recommendations are based on the general principles presented in Article 4.1.1. The recommendations outline for Members the basic elements that need to be taken into account in the design and implementation of an animal identification system to achieve animal traceability. Whatever animal identification system the country adopts, it should comply with relevant OIE standards, including Chapters 5.10. to 5.12. for animals and animal products intended for export. Each country should design a programme in accordance with the scope and relevant performance criteria to ensure that the desired animal traceability outcomes can be achieved.

Article 4.2.2.

Glossary

For the purpose of this Chapter:

Desired outcomes: describe the overall goals of a programme and are usually expressed in qualitative terms, e.g. ‘to help ensure that animals and/or animal products are safe and suitable for use’. Safety and suitability for use could be defined in terms such as animal health, food safety, trade and aspects of animal husbandry.

Performance criteria: are specifications for performance of a programme and are usually expressed in quantitative terms, such as ‘all animals can be traced to the establishment of birth within 48 hours of an enquiry’.

Reporting: means advising the Veterinary Authority and other partner organisations as appropriate in accordance with the procedures listed in the programme.

Scope: specifies the targeted species, population and/or production/trade sector within a defined area (country, zone) or compartment that is the subject of the identification and traceability programme.

Transhumance: periodic/seasonal movements of animals between different pastures within or between countries.

Article 4.2.3.

Key elements of the animal identification system

1. Desired outcomes

Desired outcomes should be defined through consultation between the Veterinary Authority and other parties, which should include (depending on scope) animal producers and food processors, private sector veterinarians, scientific research organisations and other government agencies. Desired outcomes may be defined in terms of any or all of the following:
Annex VI (contd)

a) animal health (e.g. disease surveillance and notification; detection and control of disease; vaccination programmes);

b) public health (e.g. surveillance and control of zoonotic diseases and food safety);

c) management of emergencies e.g. natural catastrophies or man-made events;

d) trade (support for inspection and certification activities of Veterinary Services, as described in Chapters 5.10. to 5.12. which reproduce model international veterinary certificates);

e) aspects of animal husbandry such as animal performance, and genetic data.

2. Scope

Scope should also be defined through consultation between the Veterinary Authority and other parties, as discussed above. The scope of animal identification systems is often based on the definition of a species and sector, to take account of particular characteristics of the farming systems e.g. pigs in pork export production; poultry in a defined compartment; cattle within a defined FMD free zone. Different systems will be appropriate according to the production systems used in countries and the nature of their industries and trade.

3. Performance criteria

Performance criteria are also designed in consultation with other parties, as discussed above. The performance criteria depend on the desired outcomes and scope of the programme. They are usually described in quantitative terms according to the epidemiology of the disease. For example, some countries consider it necessary to trace susceptible animals within 24-48 hours when dealing with highly contagious diseases such as FMD and avian influenza. For food safety, animal tracing to support investigation of incidents may also be urgent. For chronic animal diseases that are not zoonoses, it may be considered appropriate that animals can be traced over a longer period.

4. Preliminary studies

In designing animal identification systems it is useful to conduct preliminary studies, which should take into account:

a) animal populations, species, distribution, herd management,

b) farming and industry structures, production and location,

c) animal health,

d) public health,

e) trade issues,

f) aspects of animal husbandry,

g) zoning and compartmentalisation,

h) animal movement patterns (including transhumance),

i) information management and communication,

j) availability of resources (human and financial),
k) social and cultural aspects,
l) stakeholder knowledge of the issues and expectations,
m) gaps between current enabling legislation and what is needed long term,
n) international experience,
o) national experience,
p) available technology options,
q) existing identification system(s),
r) expected benefits from the animal identification systems and animal traceability and to whom they accrue,
s) issues pertaining to data ownership, access rights,
t) reporting requirements.

Pilot projects may form part of the preliminary study to test the animal identification system and animal traceability and to gather information for the design and the implementation of the programme. Economic analysis may consider costs, benefits, funding mechanisms and sustainability.

5. Design of the programme

a) General provisions

The programme should be designed in consultation with the stakeholders to facilitate the implementation of the animal identification system and animal traceability. It should take into account the scope, performance criteria and desired outcomes as well as the results of any preliminary study.

All the specified documentation should be standardised as to format, content and context.

To protect and enhance the integrity of the system, procedures should be incorporated into the design of the programme to prevent, detect and correct errors e.g. use of algorithms to prevent duplication of identification numbers and to ensure plausibility of data.

b) Means of animal identification

The choice of a physical animal or group identifier should consider elements such as the durability, human resources, species and age of the animals to be identified, required period of identification, cultural aspects, animal welfare, technology, compatibility and relevant standards, farming practices, production systems, animal population, climatic conditions, resistance to tampering, trade considerations, cost, and retention and readability of the identification method.

The Veterinary Authority is responsible for approving the materials and equipment chosen, to ensure that these means of animal identification comply with technical and field performance specifications, and for the supervision of their distribution. The Veterinary Authority is also responsible for ensuring that identifiers are unique and are used in accordance with the requirements of the animal identification system.
Annex VI (contd)

The Veterinary Authority should establish procedures for animal identification and animal traceability including:

i) the time period within which an animal born on an establishment should be identified;

ii) when animals are introduced into an establishment;

iii) when an animal loses its identification or the identifier becomes unusable;

iv) arrangements and rules for the destruction and/or reuse of identifiers;

v) penalties for the tampering and/or removal of official animal identification devices.

Where group identification without a physical identifier is adequate, documentation should be created specifying at least the number of animals in the group, the species, the date of identification, the person legally responsible for the animals and/or establishment. This documentation constitutes a unique group identifier and it should be updated to be traceable if there are any changes.

Where all animals in the group are physically identified with a group identifier, documentation should also specify the unique group identifier.

c) Registration

Procedures need to be incorporated into the design of the programme in order to ensure that relevant events and information are registered in a timely and accurate manner.

Depending on the scope, performance criteria and desired outcomes, records as described below should specify, at least, the species, the unique animal or group identifier, the date of the event, the identifier of the establishment where the event took place, and the code for the event itself.

i) Establishments/ owners or responsible keepers

Establishments where animals are kept should be identified and registered, including at least their physical location (such as geographical coordinates or street address), the type of establishment and the species kept. The register should include the name of the person legally responsible for the animals at the establishment.

The types of establishments that may need to be registered include holdings (farms), assembly centres (e.g. agriculture shows and fairs, sporting events, transit centres, breeding centres), markets, abattoirs, rendering plants, dead stock collection points, transhumance areas, centres for necropsy and diagnosis, research centres, zoos, border posts, quarantine stations.

In cases where the registration of establishments is not applicable e.g. some transhumance systems, the animal owner, the owner’s place of residence and the species kept should be recorded.

ii) Animals

Animal identification and species should be registered for each establishment/ owner. Other relevant information about the animals at each establishment/ owner may also be recorded e.g. date of birth, production category, sex, breed, animal identification of the parents.
iii) Movements

The registration of animal movements is necessary to achieve animal traceability. When an animal is introduced into or leaves an establishment, these events constitute a movement.

Some countries classify birth, slaughter and death of the animal as movements.

The information registered should include the date of the movement, the establishment from which the animal or group of animals was dispatched, the number of animals moved, the destination establishment, and any establishment used in transit.

When establishments are not registered as part of the animal identification system, ownership and location changes constitute a movement record. Movement recording may also include means of transport and the vehicle/identifier.

Procedures should be in place to maintain animal traceability during transport and when animals arrive at and leave an establishment.

iv) Events other than movements

The following events may also be registered:

- birth, slaughter and death of the animal (when not classified as a movement),
- attachment of the unique identifier to an animal,
- change of ownership or keeper regardless of change of establishment,
- observation of an animal on an establishment (testing, health investigation, health certification, etc.),
- animal imported: a record of the animal identification from the exporting country should be kept and linked with the animal identification assigned in the importing country,
- animal exported: a record of the animal identification from the exporting country should be provided to the Veterinary Authority in the importing country,
- animal identifier lost or replaced,
- animal missing (lost, stolen, etc.),
- animal identifier retired (at slaughter, following loss of the identifier or death of the animal on a farm, at diagnostic laboratories, etc.).

d) Documentation

Documentation requirements should be clearly defined and standardised, according to the scope, performance criteria and desired outcomes and supported by the legal framework.

e) Reporting

Depending on the scope, performance criteria and desired outcomes, relevant information (such as animal identification, movement, events, changes in numbers of livestock, establishments) should be reported to the Veterinary Authority by the person responsible for the animals.
f) Information system

An information system should be designed according to the scope, performance criteria and desired outcomes. This may be paper based or electronic. The system should provide for the collection, compilation, storage and retrieval of information on matters relevant to registration. The following considerations are important:

- have the potential for linkage to traceability in the other parts of the food chain;
- minimize duplication;
- relevant components, including databases, should be compatible;
- confidentiality of data;
- appropriate safeguards to prevent the loss of data, including a system for backing up the data.

The Veterinary Authority should have access to this information system as appropriate to meet the scope, performance criteria and desired outcomes.

g) Laboratories

The results of diagnostic tests should record the animal identifier or the group identifier and the establishment where the sample was collected.

h) Abattoirs, rendering plants, dead stock collection points, markets and assembly centres

Abattoirs, rendering plants, dead stock collection points, markets and assembly centres should document arrangements for the maintenance of animal identification and animal traceability in compliance with the legal framework.

These establishments are critical points for control of animal health and food safety.

Animal identification should be recorded on documents accompanying samples collected for analysis.

The components of the animal identification system operating within abattoirs should complement and be compatible with arrangements for tracking animal products throughout the food chain. At an abattoir, animal identification should be maintained during the processing of the animal’s carcass until the carcass is deemed fit for human consumption.

The animal identification and the establishment from which the animal was dispatched should be registered by the abattoir, rendering plant and dead stock collection points.

Abattoirs, rendering plants and dead stock collection points should ensure that identifiers are collected and disposed of according to the procedures established and regulated within the legal framework. These procedures should minimize the risk of unauthorized reuse and, if appropriate, should establish arrangements and rules for the reuse of identifiers.

Reporting of movement by abattoirs, rendering plants and dead stock collection points should occur according to the scope, performance criteria and desired outcomes and the legal framework.
i) **Penalties**

Different levels and types of penalties should be defined in the programme and supported by the legal framework.

6. **Legal framework**

The Veterinary Authority, with other relevant governmental agencies and in consultation with stakeholders, should establish a legal framework for the implementation and enforcement of animal identification system and animal traceability in the country. The structure of this framework will vary from country to country.

Animal identification, animal traceability and animal movement should be under the responsibility of the Veterinary Authority.

This legal framework should address:

i) desired outcomes and scope;

ii) obligations of the Veterinary Authority and other parties;

iii) organisational arrangements, including the choice of technologies and methods used for the animal identification system and animal traceability;

iv) management of animal movement;

v) confidentiality of data;

vi) data access / accessibility;

vii) checking, verification, inspection and penalties;

viii) where relevant, funding mechanisms;

ix) where relevant, arrangements to support a pilot project.

7. **Implementation**

a) **Action plan**

For implementing the animal identification system, an action plan should be prepared specifying the timetable and including the milestones and performance indicators, the human and financial resources, and checking, enforcement and verification arrangements.

The following activities should be addressed in the action plan:

i) **Communication**

The scope, performance criteria, desired outcomes, responsibilities, movement and registration requirements and sanctions need to be communicated to all parties.

Communication strategies need to be targeted to the audience, taking into account elements such as the level of literacy (including technology literacy) and spoken languages.
Annex VI (contd)

ii) Training programmes

It is desirable to implement training programmes to assist the Veterinary Services and other parties.

iii) Technical support

Technical support should be provided to address practical problems.

b) Checking and verification

Checking activities should start at the beginning of the implementation to detect, prevent and correct errors and to provide feedback on programme design.

Verification should begin after a preliminary period as determined by the Veterinary Authority in order to determine compliance with the legal framework and operational requirements.

c) Auditing

Auditing should be carried out under the authority of the Veterinary Authority to detect any problems with the animal identification system and animal traceability and to identify possible improvements.

d) Review

The programme should be subject to periodic review, taking into account the results of checking, verification and auditing activities.

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CHAPTER 4.3.

ZONING AND COMPARTMENTALISATION

Article 4.3.1.

Introduction

For the purposes of the Terrestrial Code, ‘zoning’ and ‘regionalisation’ have the same meaning.

Establishing and maintaining a disease free-status throughout the country should be the final target goal for OIE Members. However, given the difficulty of establishing and maintaining a disease free status for an entire territory, especially for diseases the entry of which is difficult to control through measures at national boundaries, there may be benefits to a Member in establishing and maintaining a subpopulation with a distinct health status within its territory. 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 Member under the provisions of this Chapter with a view to defining subpopulations of distinct health status within its territory for the purpose of disease control and/or international trade. While zoning applies to an animal subpopulation defined primarily on a geographical basis (using natural, artificial or legal boundaries), compartmentalisation applies to an animal subpopulation defined primarily by management and husbandry practices related to biosecurity. In practice, spatial considerations and good management including biosecurity plans play important roles in the application of both concepts.

A particular application of the concept of zoning is the establishment of a containment zone. In the event of a limited outbreak of a specified disease within an otherwise free country or zone, a single containment zone, which includes all cases, can be established for the purpose of minimizing the impact on the entire country or zone.

This Chapter is to assist OIE Members wishing to establish and maintain different subpopulations within their territory 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 disease outbreaks.

Before trade in animals or their products may occur, an importing country needs to be satisfied that its 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.

As well as contributing to the safety of international trade, zoning and compartmentalisation may assist disease control or eradication within a Member’s territory. Zoning may encourage the more efficient use of resources within certain parts of a country and compartmentalisation may allow the functional separation of a subpopulation from other domestic or wild animals through biosecurity measures, which a zone (through geographical separation) would not achieve. Following a disease outbreak, the use of compartmentalisation may allow a Member 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 continuation of trade.
Annex VII (contd)

Zoning and compartmentalisation cannot be applied 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 free status following a disease outbreak in a zone or compartment, Members should follow the recommendations in the relevant disease Chapter in the Terrestrial Code.

Article 4.3.2.

General considerations

The Veterinary Services of an exporting country which is establishing a zone or compartment within its territory for international trade purposes should clearly define the subpopulation in accordance with the recommendations in the relevant Chapters in the Terrestrial Code, including those on surveillance, and the identification and traceability of live animals. The Veterinary Services of an exporting country should be able to explain to the Veterinary Services of an importing country the basis for claiming a distinct animal health status for the given zone or compartment under consideration.

The procedures used to establish and maintain the distinct 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 and applicable biosecurity measures.

The authority, organisation and infrastructure of the Veterinary Services, including laboratories, must be clearly documented in accordance with the Chapter on the evaluation of Veterinary Services of the Terrestrial Code, to provide confidence in the integrity of the zone or compartment. The final authority of the zone or compartment, for the purposes of domestic and international trade, lies with the Veterinary Authority.

In the context of maintaining the health status of a population, references to ‘import’, ‘importation’ and ‘imported animals/products’ found in the Terrestrial Code apply both to importation into a country and to the movement of animals and their products into zones and compartments. Such movements should be the subject of appropriate measures to preserve the animal health status of the zone or compartment.

The exporting country should be able to demonstrate, through detailed documentation provided to the importing country, that it has implemented the recommendations in the Terrestrial 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 Terrestrial Code are applied and the Veterinary Authority of the exporting country certifies that this is the case.

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 Veterinary Services (and of the relevant industry, in the case of a compartment) including disease surveillance and diagnosis.

Biosecurity and surveillance are essential components of zoning and compartmentalisation, and the arrangements should be developed through cooperation of industry and Veterinary Services.

Industry’s responsibilities include the application of biosecurity measures, documenting and recording movements of animals and personnel, quality assurance schemes, monitoring the efficacy of the measures, documenting corrective actions, conducting surveillance, rapid reporting and maintenance of records in a readily accessible form.
The Veterinary Services should provide movement certification, and carry out documented periodic inspections of facilities, biosecurity measures, records and surveillance procedures. Veterinary Services should conduct or audit surveillance, reporting and laboratory diagnostic examinations.

Article 4.3.3.

Principles for defining a zone or compartment, including containment zone

In conjunction with the above considerations, the following principles should apply when Members define a zone or a compartment.

1. The extent of a zone and its geographical limits should be established by the Veterinary Authority on the basis of natural, artificial and/or legal boundaries, and made public through official channels.

2. In the event of an outbreak in a country or zone previously free of a disease, a containment zone may be established for the purposes of trade. Establishment of a containment zone should be based on a rapid response including:

   a) An appropriate standstill of movement of animals and commodities upon notification of suspicion of the specified disease and the demonstration that the outbreaks are contained within this zone through epidemiological investigation (trace-back, trace-forward) after confirmation of infection. The primary outbreak and likely source of the outbreak should be identified and all cases shown to be epidemiologically linked. For the effective establishment of a containment zone, it is necessary to demonstrate that there have been no new cases in the containment zone within a minimum of two incubation periods from the last detected case.

   b) A stamping-out policy or another effective control strategy aimed at eradicating the disease should be applied and the susceptible animal population within the containment zones should be clearly identifiable as belonging to the containment zone. Increased passive and targeted surveillance in accordance with Chapter 8.5.1.4 in the rest of the country or zone should be carried out and has not detected any evidence of infection.

   c) Measures consistent with the disease specific Chapter should be in place to prevent spread of the infection from the containment zone to the rest of the country or zone, including ongoing surveillance in the containment zone.

   d) For the effective establishment of a containment zone, it is necessary to demonstrate that there have been no new cases in the containment zone within a minimum of two incubation periods from the last detected case.

   e) The free status of the areas outside the containment zone would be suspended pending the establishment of the containment zone. The suspension of free status of these areas could be lifted if the containment zone is clearly established, irrespective of the provisions of the disease specific Chapter.

   f) The containment zone should be managed in such a way that it can be demonstrated that commodities for international trade can be shown to have originated outside the containment zone.

   g) The recovery of the free status of the containment zone should follow the provisions of the disease specific Chapter.
Annex VII (contd)

3. The factors defining a compartment should be established by the Veterinary Authority on the basis of relevant criteria such as management and husbandry practices related to biosecurity, and made public through official channels. Animals and herds belonging to such subpopulations need to be recognisable as such through a clear epidemiological separation from other animals and all things presenting a disease risk.

4. For a zone or compartment, the Veterinary Authority should document in detail the measures taken to ensure the identification of the subpopulation and the establishment and maintenance of its health status through a biosecurity plan. The measures used to establish and maintain the distinct 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 health status of animals in adjacent areas, applicable biosecurity measures (including movement controls, use of natural and artificial boundaries, the spatial separation of animals, and commercial management and husbandry practices), and surveillance.

5. Relevant animals within the zone or compartment should be identified in such a way that their history can be audited. Depending on the system of production, identification may be done at the herd, flock, lot or individual animal level. Relevant animal movements into and out of the zone or compartment should be well documented, controlled and supervised. The existence of a valid animal identification system is a prerequisite to assess the integrity of the zone or compartment.

6. For a compartment, the biosecurity plan should describe the partnership between the relevant industry and the Veterinary Authority, and their respective responsibilities. It should also describe the routine operating procedures to provide clear evidence that the surveillance conducted, the live animal identification and traceability system, and the management practices are adequate to meet the definition of the compartment. In addition to information on animal movement controls, the plan should include herd or flock production records, feed sources, surveillance results, birth and death records, visitor logbook, morbidity and mortality history, medications, vaccinations, documentation of training of relevant personnel and any other criteria necessary for evaluation of risk mitigation. The information required may vary according to the 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.

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CHAPTER 4.4.
APPLICATION OF COMPARTMENTALISATION

Article 4.4.1.

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 4.3. with the objective to facilitate trade in animals and products of animal origin and as a tool for disease management.

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

The Glossary of the Terrestrial Code defines a compartment as “an animal subpopulation contained in one or more establishments under a common biosecurity management system with a distinct health status with respect to a specific disease or specific diseases for which required surveillance, control and biosecurity measures have been applied for the purpose of international trade”.

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 practices and biosecurity. However, spatial considerations and good management practices play a role in the application of both concepts.

Compartmentalisation is not a new concept for Veterinary Services; in fact, it has been applied for a long time in many disease control programmes that are based on the concept of disease-free herds/flocks.

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 animal production operation in an infected country or zone might have biosecurity measures and management practices that result in negligible risk from disease 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 zones, compartments preferably should be defined prior to the occurrence of a disease outbreak. In the event of an outbreak or in infected countries or zones, compartmentalisation may be used to facilitate trade.

For the purpose of international trade, compartments must be under the responsibility of the Veterinary Authority in the country. For the purposes of this Chapter, compliance by the Members with Chapters 1.1. and 3.1. is an essential prerequisite.
Annex VII (contd)

Article 4.4.2.

**Principles for defining a compartment**

A compartment may be established with respect of a specific disease or diseases. A compartment must be clearly defined, indicating the location of all its components including establishments as well as related functional units (such as feed mills, slaughterhouses, rendering plants, etc.), their interrelationships and their contribution to an epidemiological separation between the animals in a compartment and subpopulations with a different health status. The definition of compartment may revolve around disease specific epidemiological factors, animal production systems, biosecurity practices infrastructural factors and surveillance.

Article 4.4.3.

**Separation of a compartment from potential sources of infection**

The management of a compartment must provide to the Veterinary Authority 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) flocks or herds with a different health status in close proximity to the compartment, including wildlife and their migratory routes;
      
      ii) slaughterhouses, rendering plants or feed mills;
      
      iii) markets, fairs, agricultural shows, sporting events, zoos, circuses and other points of animal concentration.

2. **Infrastructural factors**

   Structural aspects of the establishments within a compartment contribute to the effectiveness of its Biosecurity. Consideration should be given to:

   a) fencing or other effective means of physical separation;
   
   b) facilities for people entry including access control, changing area and showers;
   
   c) vehicle access including washing and disinfection procedures;
   
   d) unloading and loading facilities;
e) isolation facilities for introduced animals;

f) facilities for the introduction of material and equipment;

g) infrastructure to store feed and veterinary products;

h) disposal of carcasses, manure and waste;

i) water supply;

j) measures to prevent exposure to living mechanical or biological vectors such as insects, rodents and wild birds;

k) air supply;

l) feed supply/source.

More detailed recommendations for certain establishments can be found in Sections 4 and 6 of the Terrestrial Code.

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 animal movements, rodents, fauna, aerosols, arthropods, 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 the 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 Veterinary 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 regularly 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 disease agent into the compartment.

4. Traceability system

A prerequisite for assessing the integrity of a compartment is the existence of a valid traceability system. All animals within a compartment should be individually identified and registered in such a way that their history and movements can be documented and audited. In cases where individual identification may not be feasible, such as broilers and day-old chicks, the Veterinary Authority should provide sufficient assurance of traceability.

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

Article 4.4.4.

Documentation

Documentation must 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 herd or flock production records, feed sources, laboratory tests, birth and death records, the visitor logbook, morbidity history, 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 Terrestrial Code Chapter.

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

Vaccination records including the type of vaccine and frequency of administration must be available to enable interpretation of surveillance data.

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

All relevant information must be recorded in a transparent manner and be easily accessible so as to be auditable by the Veterinary Authority.
Surveillance for the 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 detection level of the internal and external surveillance should be reviewed and, where necessary, raised. 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 Veterinary 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 must 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 active 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.

Diagnostic capabilities and procedures

Officially-designated laboratory facilities complying with the OIE standards for quality assurance, as defined in Chapter 1.1.2. of the Terrestrial Manual, should be available for sample testing. All laboratory tests and procedures should comply with the recommendations of the Terrestrial Manual for the specific disease.

Each laboratory that conducts testing should have systematic procedures in place for rapid reporting of disease results to the Veterinary Authority. Where appropriate, results should be confirmed by an OIE Reference Laboratory.

Emergency response and notification

Early detection, diagnosis and notification of disease are critical to minimize the consequences of outbreaks.
Annex VII (contd)

In the event of suspicion of occurrence of the disease for which the compartment was defined, export certification 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 an occurrence of any infectious disease not present according to the baseline animal health report of the compartment referred to in Article 4.4.4., the management of the compartment should notify the Veterinary Authority, and initiate a review to determine whether there has been a breach in the biosecurity measures. If a significant breach in biosecurity, even in the absence of outbreak, was detected, export certification of the compartment should be suspended. Trade Disease-Free status of the compartment may only be resumed after the compartment has adopted the necessary measures to re-establish the original biosecurity level and the Veterinary Authority re-approves the status of the compartment.

In the event of a compartment being in close proximity to an outbreak of the at risk from a change in the surrounding area, in the disease situation for which the compartment was defined, the Veterinary Authority should re-evaluate without delay the status of the compartment and consider additional biosecurity measures applied to ensure that the integrity of the compartment has been maintained.

Article 4.4.8.

Supervision and control of a compartment

The authority, organisation, and infrastructure of the Veterinary Services, including laboratories, must be clearly documented in accordance with the Chapter on the Evaluation of Veterinary Services of the Terrestrial Code, to provide confidence in the integrity of the compartment.

The Veterinary Authority has the final authority in granting, suspending and revoking the status of a compartment. The Veterinary 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.

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CHAPTER 3.X.X.

GENERAL GUIDELINES FOR SURVEILLANCE OF ARTHROPOD VECTORS OF ANIMAL DISEASES

Article 1

1. Introduction

Vector-borne diseases are of increasing importance economically and to human and animal health.

Environmental (including climate change), sociological and economical changes may affect the distribution and impact of these diseases.

Improved understanding of the distribution and population dynamics of the vectors is a key element for assessing and managing the risks associated with vector-borne animal and zoonotic diseases.

The Terrestrial Code contains recommendations for the surveillance of several vector-borne diseases.

The need has arisen to complement these general recommendations on surveillance guidelines with additional general guidelines advice on for the surveillance of vectors themselves. This Chapter only addresses surveillance for arthropod vectors.

For the purpose of trade, it must be noted that there is no conclusive relationship between the presence of a vector(s) and the disease status of a country/zone and also that the apparent absence of a vector(s) does not by itself confirm vector-free status.

A vector may be broadly defined as:

"...in infectious disease epidemiology, an insect or any living carrier that transports an infectious agent from an infected individual or its wastes to a susceptible individual or its food or immediate surroundings. The organism may or may not pass through a development cycle within the vector". (Dictionary of Epidemiology, John M. last, 4th Edition 2001)

A decision tree with respect to vector surveillance is reflected presented in Figure 1.
2. Objectives

The objective of these recommendations is to provide methods for:

- gathering up-to-date information on the spatial and temporal distribution and abundance of vectors of the arthropod-borne OIE-listed diseases and emerging diseases;
- monitoring changes in the spatial and temporal distribution and abundance of these vectors;
- collecting relevant data to inform risk assessment (including vector competency) and risk management of these vector-borne diseases;
- detecting the presence of specific vectors or confirming their absence;
- understanding pathways of entry for vectors and vector-borne pathogenic agents.
Article 3.

3. Sampling methodology

a) Sampling plan

i) The state of existing knowledge should be assessed to determine whether or not historical data on the vector or the disease exist for the country or zone. The objective of the surveillance programme should be determined and stated before planning begins.

ii) All available historical data on the vector or the disease for the country or zone should be collated and assessed.

iii) The sampling plan should consider the following:

- the known aspects of the biology and ecology of the vector(s),
- the presence, distribution and abundance of the vectors’ host animal population(s),
- the environmental, climatic, ecological and topographic conditions of relevance to vector ecology,
- the need for a risk assessment to indicate the areas at highest risk of introduction of a vector that is unlikely to be present.

iv) Sampling should be aimed at:

- establishing vector presence or confirming vector absence in the country or zone,
- describing the distribution of the vector(s) within the country or zone,
- providing additional information on vector density and spatial/temporal variability (both over the short- and the long-term),
- early detection of vectors or vector-borne pathogenic agents in areas with risk of entry and establishment.

iv) The sampling plan should be designed to provide representative appropriate estimates, with a minimum of bias, of the indicators listed in item 3 above. Consideration should be given to the following:

The recommended general approach to sampling is via a three-stage hierarchy:

- Stratification based on ecological criteria (where possible), and risk assessment for vector introduction,
- subdivision of strata into spatial sampling units, and
- establishment of actual sampling sites within selected spatial sampling units.
Annex VIII (contd)

If adequate epidemiological and historical data and/or expert opinion are available, the sampling plan may be further refined or targeted by defining strata which are as homogenous as possible with respect to the following known or suspected risk-factors, as appropriate for the country or zone:

- domestic or wild populations of host animals preferred by the vector,
- vector habitat suitability,
- climatic patterns (including seasonal),
- areas endemically and/or epidemically affected by the disease of concern,
- areas of known vector occurrences,
- fringe zone(s) around areas of known vector occurrences or other high risk areas for vector introduction, such as port,
- areas in which the disease(s) or vector(s) of concern have not been reported currently or historically,
- each stratum (or the whole country or zone if not stratified) should be divided into spatial sampling units according to standard methodologies such as a grid system,
- the number and size of the spatial sampling units should be defined to provide representative appropriate estimates of the indicators listed in item 3 above,
- the number and location of actual sampling sites within each spatial sampling unit also should be defined to provide representative appropriate estimates of the indicators listed in item 3 above,
- different levels of sampling intensity (spatial sampling unit size, number of units sampled, number of sites sampled within units, and sampling frequency) may be applied to different strata into which the country or zone has been divided. For example, more intensive sampling might be carried out in strata where vector presence seems most likely, based on biological or statistical criteria.

b) Sampling methods

Many sampling methods have been developed for the capture of vector arthropods, and these differ according to the disease/vector system under consideration.

i) The collection methods used should be adapted as required to ensure reasonable confidence of collecting the vector(s) of concern.

ii) Collection methods should secure obtain the various developmental stages (such as eggs, larvae, nymphs, adults) and adult age categories, as appropriate to the species in question, required to estimate population survival rates and population dynamics in relation to disease transmission.
iii) Different collection methods may be required to secure obtain samples from a single vector species, depending on the life stage or place of capture (such as from the environment or from the host animals). The collection method must be appropriate to the species and life stage of interest.

The collection methods should preserve the vector(s) to allow for subsequent complete taxonomic analysis (using both morphological and molecular techniques) and detection and/or isolation of pathogenic agent(s), in a manner suitable for their morphological identification or identification with molecular techniques. Where the purpose of sampling is to detect or isolate a pathogenic agent(s), specific protocols should be followed to ensure the samples are suitable for these assays.

c) Data management, analysis and interpretation

Data management and analytical methodologies should be done in accordance with Chapter 1.4.).
CHAPTER 4.5.

COLLECTION AND PROCESSING OF BOVINE, SMALL RUMINANT AND PORCINE SEMEN

Article 4.5.1.

General considerations

The purposes of official sanitary control of semen production are to:

1. maintain the health of animals on an artificial insemination centre at a level which permits the international distribution of semen with a negligible risk of infecting other animals or humans with pathogens transmissible by semen;

2. ensure that semen is hygienically collected, processed and stored.

Artificial insemination centres should comply with recommendations in Chapter 4.6.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 4.5.2.

Conditions applicable to testing of bulls and teaser animals

Bulls and teaser animals should only enter an artificial insemination centre if they fulfil the following requirements.

1. Pre-quarantine

The animals should comply with the following requirements prior to entry into isolation at the quarantine station.

a) Bovine brucellosis

The animals should comply with point 3 or 4 of Article 11.3.5.

b) Bovine tuberculosis

The animals should comply with point 3 or 4 of Article 11.7.5.

c) Bovine viral diarrhoea-mucosal disease (BVD-MD)

The animals should be subjected to the following tests:

i) a virus isolation test or a test for virus antigen, with negative results;

ii) a serological test to determine the serological status of every animal.
Annex IX (contd)

d) Infectious bovine rhinotracheitis-infectious pustular vulvovaginitis

If the artificial insemination centre is to be considered as infectious bovine rhinotracheitis-infectious pustular vulvovaginitis free (IBR/IPV), the animals should either:

i) come from an IBR/IPV free herd as defined in Article 11.12.3.; or

ii) be subjected, with negative results, to a serological test for IBR/IPV on a blood sample.

e) Bluetongue

The animals should comply with Articles 8.3.5., 8.3.6. or 8.3.7., depending on the bluetongue status of the country of origin of the animals.

2. Testing in the quarantine station prior to entering the semen collection facilities

Prior to entering the semen collection facilities of the artificial insemination centre, bulls and teaser animals should be kept in a quarantine station for at least 28 days. The animals should be subjected to diagnostic tests as described below a minimum of 21 days after entering the quarantine station, except for Campylobacter fetus subsp. venerealis and Trimatophomonas fetus, for which testing may commence after 7 days in quarantine. All the results should be negative except in the case of BVD-MD antibody serological testing (see point 2b)i) below).

a) Bovine brucellosis

The animals should be subjected to a serological test with negative results.

b) BVD-MD

i) All animals should be tested for viraemia as described in point 1c) above.

Only when all the animals in quarantine test negative for viraemia, may the animals enter the semen collection facilities upon completion of the 28-day quarantine period.

ii) After 21 days in quarantine, all animals should be subjected to a serological test to determine the presence or absence of BVD-MD antibodies.

iii) Only if no sero-conversion occurs in the animals which tested seronegative before entry into the quarantine station, may any animal (seronegative or seropositive) be allowed entry into the semen collection facilities.

iv) If sero-conversion occurs, all the animals that remain seronegative should be kept in quarantine over a prolonged time until there is no more seroconversion in the group for a period of 3 weeks. Serologically positive animals may be allowed entry into the semen collection facilities.

c) Campylobacter fetus subsp. venerealis

i) Animals less than 6 months old or kept since that age only in a single sex group prior to quarantine should be tested once on a preputial specimen, with a negative result.

ii) Animals aged 6 months or older that could have had contact with females prior to quarantine should be tested three times at weekly intervals on a preputial specimen, with a negative result in each case.
d) *Tritrichomonas foetus*

i) Animals less than 6 months old or kept since that age only in a single sex group prior to quarantine, should be tested once on a preputial specimen, with a negative result.

ii) Animals aged 6 months or older that could have had contact with females prior to quarantine should be tested three times at weekly intervals on a preputial specimen, with a negative result in each case.

e) **IBR-IPV**

If the artificial insemination centre is to be considered as IBR/IPV free, the animals should be subjected, with negative results, to a diagnostic test for IBR/IPV on a blood sample. If any animal tests positive, the animal should be removed immediately from the quarantine station and the other animals of the same group should remain in quarantine and be retested, with negative results, not less than 21 days after removal of the positive animal.

f) **Bluetongue**

The animals should comply with Articles 8.3.5., 8.3.6. or 8.3.7., depending on the bluetongue status of the country of origin of the animals.

3. **Testing for BVD-MD prior to the initial dispatch of semen from each serologically positive bull**

Prior to the initial dispatch of semen from BVD-MD serologically positive bulls, a semen sample from each animal should be subjected to a virus isolation or virus antigen test for BVD-MD. In the event of a positive result, the bull should be removed from the centre and all of its semen destroyed.

4. **Testing of frozen semen for IBR/IPV in artificial insemination centres not considered as IBR/IPV free**

Each aliquot of frozen semen should be tested as per Article 11.12.7.

5. **Testing programme for bulls and teasers resident in the semen collection facilities**

All bulls and teasers resident in the semen collection facilities should be tested at least annually for the following diseases, with negative results, where the country of origin is not free:

a) Bovine brucellosis

b) Bovine tuberculosis

c) **BVD-MD**

A nimals negative to previous serological tests should be retested to confirm absence of antibodies. Should an animal become serologically positive, every ejaculate of that animal collected since the last negative test should be either discarded or tested for virus with negative results.

d) **Campylobacter fetus subsp. venerealis**

i) A preputial specimen should be cultured.
Annex IX (contd)

ii) Only bulls on semen production or having contact with bulls on semen production need to be tested. Bulls returning to collection after a lay off of more than 6 months should be tested not more than 30 days prior to resuming production.

e) Bluetongue

The animals should comply with the provisions referred to in Articles 8.3.5., 8.3.6. or 8.3.7., depending on the bluetongue status of the country of origin of the animals.

f) Trichomonas foetus

i) A preputial specimen should be cultured.

ii) Only bulls on semen production or having contact with bulls on semen production need to be tested. Bulls returning to collection after a lay off of more than 6 months should be tested not more than 30 days prior to resuming production.

g) IBR-IPV

If the artificial insemination centre is to be considered as IBR/IPV free, the animals should comply with the provisions in point 2)(c) of Article 11.12.3.

Article 4.5.3.

Conditions applicable to testing of rams/ bucks and teaser animals

Rams/ bucks and teaser animals should only enter an artificial insemination centre if they fulfil the following requirements.

1. Pre-quarantine

The animals should comply with the following requirements prior to entry into isolation at the quarantine station.

a) Caprine and ovine brucellosis

The animals should comply with Article 14.1.6.

b) Ovine epididymitis

The animals should comply with Article 14.7.3.

c) Contagious agalactia

The animals should comply with points 1 and 2 of Article 14.3.1.

d) Peste des petits ruminants

The animals should comply with points 1, 2, and 4 or 5 of Article 14.8.7.

e) Contagious caprine pleuropneumonia

The animals should comply with Article 14.4.5. or Article 14.4.7., depending on the contagious caprine pleuropneumonia status of the country of origin of the animals.
f) Paratuberculosis

The animals should be free from clinical signs for the past 2 years.

g) Scrapie

If the animals do not originate from a scrapie free country or zone as defined in Article 14.9.3., the animals should comply with Article 14.9.6.

h) Maedi-visna

The animals should comply with Article 14.6.2.

i) Caprine arthritis/encephalitis

In the case of goats, the animals should comply with Article 14.2.2.

j) Bluetongue

The animals should comply with Articles 8.3.5., 8.3.6. or 8.3.7., depending on the bluetongue status of the country of origin of the animals.

k) Tuberculosis

In the case of goats, the animals should be subject to a single or comparative tuberculin test, with negative results.

l) Border disease

The animals should be subject to a viral agent isolation test with negative results.

2. Testing in the quarantine station prior to entering the semen collection facilities

Prior to entering the semen collection facilities of the artificial insemination centre, rams/bucks and teasers should be kept in a quarantine station for at least 28 days. The animals should be subjected to diagnostic tests as described below a minimum of 21 days after entering the quarantine station, with negative results.

a) Caprine and ovine brucellosis

The animals should be subject to testing as described in point 1c) of Article 14.1.8.

b) Ovine epididymitis

The animals and semen should be subject to testing as described in points 1d) and 2 of Article 14.7.4.

c) Maedi-visna and caprine arthritis/encephalitis

The animals and semen should be subjected to a serological test.
Annex IX (contd)

d) Bluetongue

The animals should comply with the provisions referred to in Articles 8.3.5., 8.3.6. or 8.3.7., depending on the bluetongue status of the country of origin of the animals.

3. Testing programme for rams/bucks and teasers resident in the semen collection facilities

All rams/bucks and teasers resident in the semen collection facilities should be tested at least annually for the following diseases, with negative results, where the country of origin is not free:

a) caprine and ovine brucellosis;

b) ovine epididymitis;

c) Maedi-visna and caprine arthritis/encephalitis;

d) tuberculosis (for goats only);

e) bluetongue.

Article 4.5.4

Conditions applicable to testing of boars

Boars should only enter an artificial insemination centre if they fulfil the following requirements.

1. Pre-quarantine

The animals should be clinically healthy, physiologically normal and comply with the following requirements within 30 days prior to entry into isolation at the quarantine station.

a) Porcine brucellosis

The animals should comply with Article 15.4.3.

b) Foot and mouth disease

The animals should comply with Article 8.5.10., 8.5.11. or 8.5.12.

c) Aujeszky’s disease

The animals should comply with Article 8.2.8. or 8.2.9.

d) Teschovirus encephalomyelitis

The animals should comply with Article 15.6.5. or 15.6.7.

e) Transmissible gastroenteritis

The animals should comply with Article 15.7.2.

f) Swine vesicular disease

The animals should comply with Article 15.5.5. or 15.5.7.
g) African swine fever
   The animals should comply with Article 15.1.5. or 15.1.6.

h) Classical swine fever
   The animals should comply with Article 15.3.7., 15.3.8. or 15.3.9.

i) Porcine reproductive and respiratory syndrome
   The animals should be subject to the test complying with the standards in the Terrestrial Manual.

2. Testing in the quarantine station prior to entering the semen collection facilities

   Prior to entering the semen collection facilities of the artificial insemination centre, boars should be kept in a quarantine station for at least 28 days. The animals should be subjected to diagnostic tests as described below a minimum of 21 days after entering the quarantine station, with negative results.

   a) Porcine brucellosis
      The animals should comply with Article 15.4.5.

   b) Foot and mouth disease
      The animals should comply with Article 8.5.13., 8.5.14., 8.5.15. or 8.5.16.

   c) Aujeszky’s disease
      The animals should comply with Article 8.2.12., 8.2.13. or 8.2.14.

   d) Teschovirus encephalomyelitis
      The animals should comply with Article 15.6.9. or 15.6.10.

   e) Transmissible gastroenteritis
      The animals should comply with Article 15.7.4.

   f) Swine vesicular disease
      The animals should comply with Article 15.5.9. or 15.5.10.

   g) African swine fever
      The animals should comply with Article 15.1.8. or 15.1.9.

   h) Classical swine fever
      The animals should comply with Article 15.3.11., 15.3.12. or 15.3.13.

   i) Porcine reproductive and respiratory syndrome
      The animals should be subject to the test complying with the standards in the Terrestrial Manual.
Annex IX (contd)

3. Testing programme for boars resident in the semen collection facilities

All boars resident in the semen collection facilities should be tested at least annually for the following diseases, with negative results, where the compartment/zone or country is not free:

a) Porcine brucellosis

The animals should comply with Article 15.4.5.

b) Foot and mouth disease

The animals should comply with Article 8.5.13., 8.5.14., 8.5.15. or 8.5.16.

c) Aujeszky’s disease

The animals should comply with Article 8.2.12., 8.2.13. or 8.2.14. regarding testing every four months.

d) Teschovirus encephalomyelitis

The animals should comply with Article 15.6.9. or 15.6.10.

e) Transmissible gastroenteritis

The animals should comply with Article 15.7.4.

f) Swine vesicular disease

The animals should comply with Article 15.5.9. or 15.5.10.

g) African swine fever

The animals should comply with Article 15.1.8. or 15.1.9. Routine test to be applied at least every six months.

h) Classical swine fever

The animals should comply with Article 15.3.11., 15.3.12. or 15.3.13.

i) Porcine reproductive and respiratory syndrome

The animals should be subject to the test complying with the standards in the Terrestrial Manual.

Article 4.5.5.

General considerations for hygienic collection and handling of semen

Observation of the recommendations described in the Articles below will very significantly reduce the likelihood of the semen being contaminated with common bacteria which are potentially pathogenic.
Article 4.5.6.

Conditions applicable to the collection of semen

1. The floor of the mounting area should be easy to clean and to disinfect. A dusty floor should be avoided.

2. The hindquarters of the teaser, whether a dummy or a live teaser animal, should be kept clean. A dummy should be cleaned completely after each period of collection. A teaser animal should have its hindquarters cleaned carefully before each collecting session. The dummy or hindquarters of the teaser animal should be sanitized after the collection of each ejaculate. Disposable plastic covers may be used.

3. The hand of the person collecting the semen should not come into contact with the animal’s penis. Disposable gloves should be worn by the collector and changed for each collection.

4. The artificial vagina should be cleaned completely after each collection where relevant. It should be dismantled, its various parts washed, rinsed and dried, and kept protected from dust. The inside of the body of the device and the cone should be disinfected before re-assembly using approved disinfection techniques such as those involving the use of alcohol, ethylene oxide or steam. Once re-assembled, it should be kept in a cupboard which is regularly cleaned and disinfected.

5. The lubricant used should be clean. The rod used to spread the lubricant should be clean and should not be exposed to dust between successive collections.

6. The artificial vagina should not be shaken after ejaculation, otherwise lubricant and debris may pass down the cone to join the contents of the collecting tube.

7. When successive ejaculates are being collected, a new artificial vagina should be used for each mounting. The vagina should also be changed when the animal has inserted its penis without ejaculating.

8. The collecting tubes should be sterile, and either disposable or sterilised by autoclaving or heating in an oven at 180°C for at least 30 minutes. They should be kept sealed to prevent exposure to the environment while awaiting use.

9. After semen collection, the tube should be left attached to the cone and within its sleeve until it has been removed from the collection room for transfer to the laboratory.

Article 4.5.7.

Conditions applicable to the handling of semen and preparation of semen samples in the laboratory

1. Diluents:

   a) All receptacles used should have been sterilised.

   b) Buffer solutions employed in diluents prepared on the premises should be sterilized by filtration (0.22 µm) or by autoclaving (121°C for 30 minutes) or be prepared using sterile water before adding egg yolk (if applicable) or equivalent additive and antibiotics.
Annex IX (contd)

c) If the constituents of a diluent are supplied in commercially available powder form, the water used must have been distilled or demineralised, sterilized (121°C for 30 minutes or equivalent), stored correctly and allowed to cool before use.

d) Whenever milk, egg yolk or any other animal protein is used in preparing the semen diluent, the product must be free of pathogens or sterilised; milk heat-treated at 92°C for 3.5 minutes, eggs from SPF flocks when available. When egg yolk is used, it should be separated from eggs using aseptic techniques. Alternatively, commercial egg yolk prepared for human consumption or egg yolk treated by, for example, pasteurisation or irradiation to reduce bacterial contamination, may be used. Other additives must also be sterilized before use.

e) Diluent should not be stored for more than 72 hours at +5°C before use. A longer storage period is permissible for storage at -20°C. Storage vessels should be stoppered.

f) A mixture of antibiotics should be included with a bactericidal activity at least equivalent to that of the following mixtures in each ml of frozen semen: gentamicin (250 µg), tylosin (50 µg), lincomycin-spectinomycin (150/300 µg); penicillin (500 IU), streptomycin (500 µg), lincomycin-spectinomycin (150/300 µg); or amikacin (75 µg), divekacin (25 µg).

The names of the antibiotics added and their concentration should be stated in the international veterinary certificate.

2. Procedure for dilution and packing

a) The tube containing freshly collected semen should be sealed as soon as possible after collection, and kept sealed until processed.

b) After dilution and during refrigeration, the semen should also be kept in a stoppered container.

c) During the course of filling receptacles for dispatch (such as insemination straws), the receptacles and other disposable items should be used immediately after being unpacked. Materials for repeated use should be disinfected with alcohol, ethylene oxide, steam or other approved disinfection techniques.

d) If sealing powder is used, care should be taken to avoid its being contaminated.

3. Conditions applicable to the storage of semen

Semen for export should be stored separately from other genetic material not meeting these recommendations in fresh liquid nitrogen in sterilised/sanitised flasks before being exported.

Semen straws should be sealed and code marked in line with the international standards of the International Committee for Animal Recording (ICAR).3

Prior to export, semen straws or pellets should be identified and placed into new liquid nitrogen in a new or sterilised flask or container under the supervision of an Official Veterinarian. The contents of the container or flask should be verified by the Official Veterinarian prior to sealing with an official numbered seal before export and accompanied by an international veterinary certificate listing the contents and the number of the official seal.

OIE Terrestrial Animal Health Standards Commission / March 2009
4. Sperm sorting

Equipment used for sex-sorting sperm should be clean and disinfected between animals according to the manufacturer’s recommendations.

Where seminal plasma, or components thereof, is added to sorted semen prior to cryopreservation and storage, it should be derived from animals of same or better health status.

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1 The ICAR international standards on straws are contained in Recording Guidelines – Appendices to the international agreement of recording practices. Section 9, Appendix B relating to semen straw identification.

The text of this document is available at the following web site: www.icar.org
CHAPTER 4.6.

GENERAL HYGIENE IN SEMEN COLLECTION AND PROCESSING CENTRES

Article 4.6.1.

General considerations

Observation of the recommendations described in the Articles below will very significantly reduce the likelihood of the semen being contaminated with common bacteria which are potentially pathogenic.

Article 4.6.2.

Conditions applicable to artificial insemination centres

1. The artificial insemination centre is comprised of:
   a) animal accommodation areas (including one isolation facility for sick animals) and a semen collection room, these two premises hereon designated as semen collection facilities; accommodation areas should be species specific where relevant;
   b) a semen laboratory and semen storage areas;
   c) administration offices.

   A quarantine station may also be attached to the centre, provided that it is on a different location from that of those two first parts.

2. The centre should be officially approved by the Veterinary Authority.

3. The centre should be under the supervision and control of the Veterinary Services which will be responsible for regular audits, at an interval of no more than 6 months, of protocols, procedures and prescribed records on the health and welfare of the animals in the centre and on the hygienic production, storage and dispatch of semen.

4. The centre should be under the direct supervision and control of an official veterinarian.

5. Only swine associated with semen production should be permitted to enter the centre. Other species of livestock may exceptionally be resident on the centre, provided that they are kept physically apart from the swine.

6. Swine on the centre should be adequately isolated from farm livestock on adjacent land or buildings for instance by natural or artificial means.

7. The entry of visitors should be strictly controlled. Personnel at a centre should be technically competent and observe high standards of personal hygiene to preclude the introduction of pathogenic organisms. Protective clothing and footwear for use only on the centre should be provided.

8. Individual semen containers and storage rooms should be capable of being disinfected.
Conditions applicable to semen collection facilities

1. The semen collection facilities should include separate and distinct areas for accommodating resident animals, for semen collection, for feed storage, for manure storage, and for the isolation of animals suspected of being infected.

2. Only animals associated with semen production should be permitted to enter the semen collection facilities. Other species of animals may be resident at the centre, if necessary for the movement or handling of the donors and teasers or for security, but contact with the donors and teasers should be minimised. All animals resident at the semen collection facilities must meet the minimum health requirements for donors.

3. The donors and teasers should be adequately isolated to prevent the transmission of diseases from farm livestock and other animals. Measures should be in place to prevent the entry of wild animals susceptible to ruminant and swine diseases transmissible via semen.

4. Personnel at the centre should be technically competent and observe high standards of personal hygiene to preclude the introduction of pathogenic organisms. Special protective clothing and footwear for use only at the semen collection facilities should be provided and worn at all times inside.

5. Visitors to the semen collection facilities should be kept to a minimum, and visits should be subject to formal authorisation and control. Equipment for use with the livestock should be dedicated to the semen collection facilities or disinfected prior to entry. All equipment and tools brought on to the premises must be examined and treated if necessary to ensure that they cannot introduce disease.

6. Vehicles used for transport of animals to and from the semen collection facilities should not be allowed to enter the facilities.

7. The semen collection area should be cleaned daily after collection. The animals' accommodation and semen collection areas should be cleaned and disinfected at least once a year.

8. Fodder introduction and manure removal should be done in a manner which poses no significant animal health risk.

Conditions applicable to semen laboratories

1. The semen laboratory should be physically separated from the semen collection facilities, and include separate areas for artificial vagina cleaning and preparation, semen evaluation and processing, semen pre-storage and storage. Entry to the laboratory should be prohibited to unauthorised personnel.

2. The laboratory personnel should be technically competent and observe high standards of personal hygiene to preclude the introduction of pathogenic organisms during semen evaluation, processing and storage.

3. Visitors to the laboratory should be kept to a minimum, and visits should be subject to formal authorisation and control.

4. The laboratory should be constructed with materials that permit effective cleaning and disinfection.
5. The laboratory should be regularly cleaned. Work surfaces for semen evaluation and processing should be cleaned and disinfected at the end of each workday.

6. The laboratory should be treated against rodents and insects on a regular basis as needed to control these pests.

7. The storage rooms and individual semen containers should be easy to clean and disinfect.

8. Only semen collected from donors having a health status equivalent to or better than the donors at the semen collection facilities should be processed in the laboratory.

Article 4.6.5.

Conditions applicable to the management of bulls, rams, bucks and boars

The objective is to keep the animals in a satisfactory state of cleanliness, particularly of the lower thorax and abdomen.

1. Whether on pasture or housed, the animal should be kept under hygienic conditions. If housed, the litter must be kept clean and renewed as often as necessary.

2. The coat of the animal should be kept clean.

3. For bulls, the length of the tuft of hairs at the preputial orifice, which is invariably soiled, should be cut to about 2 cm. The hair should not be removed altogether, because of its protective role. If cut too short, irritation of the preputial mucosa may result because these hairs aid the drainage of urine.

4. The animal should be brushed regularly, and where necessary on the day before semen collection, paying special attention to the underside of the abdomen.

5. In the event of obvious soiling, there should be careful cleaning, with soap or a detergent, of the preputial orifice and the adjoining areas, followed by thorough rinsing and drying.

6. When the animal is brought into the collection area, the technician must make sure that it is clean, and that it is not carrying any excessive litter or particles of feed on its body or its hooves, for such materials are always heavily contaminated.
CHAPTER 4.7.

COLLECTION AND PROCESSING OF IN VIVO DERIVED EMBRYOS FROM LIVESTOCK AND HORSES

Article 4.7.1.

Aims of control

The purpose of official sanitary control of in vivo derived embryos intended for movement internationally is to ensure that specific pathogenic organisms, which could be associated with embryos, are controlled and transmission of infection to recipient animals and progeny is avoided.

Article 4.7.2.

Conditions applicable to the embryo collection team

The embryo collection team is a group of competent technicians, including at least one veterinarian, to perform the collection, processing and storage of embryos. The following conditions should apply:

1. The team should be supervised by a team veterinarian.

2. The team veterinarian is responsible for all team operations which include verification of donor health status, sanitary handling and surgery of donors, and disinfection and hygienic procedures.

3. The team veterinarian should be specifically approved for this purpose by an Official Veterinarian.

4. Team personnel should be adequately trained in the techniques and principles of disease control. High standards of hygiene should be practiced to preclude the introduction of infection.

5. The collection team must have adequate facilities and equipment for:

   a) collecting embryos;

   b) processing and treatment of embryos at a permanent site or mobile laboratory;

   c) storing embryos.

   These facilities need not necessarily be at the same location.

6. The embryo collection team must keep a record of its activities, which must be maintained for inspection by the approving Veterinary Authority for a period of at least 2 years after the embryos have been exported.

7. The embryo collection team should be subjected to regular inspection at least once a year by an Official Veterinarian to ensure compliance with procedures for the sanitary collection, processing and storage of embryos.
Annex IX (contd)

8. The collection team must not operate in an infected zone with regard to foot and mouth disease (except for the collection of in vivo derived bovine embryos), rinderpest, peste des petits ruminants, contagious bovine pleuropneumonia, African horse sickness, African swine fever and classical swine fever.

Article 4.7.3.

Conditions applicable to the processing laboratories

The processing laboratory used by the embryo collection team may be mobile or permanent. It is a facility in which embryos are recovered from collection media, examined and subjected to any required treatments such as washing before freezing, storage and quarantine, pending results of diagnostic procedures, and being examined and prepared for freezing and storage.

A permanent laboratory may be part of a specifically designed collection and processing unit, or a suitably adapted part of an existing building. It may be on the premises where the donor animals are kept. In either case, the laboratory should be physically separated from animals. Both mobile and permanent laboratories should have a clear separation between dirty areas (animal handling) and the clean processing area.

Additionally:

1. The processing laboratory should be under the direct supervision of the team veterinarian and be regularly inspected by an Official Veterinarian.

2. While embryos for export are being handled prior to their storage in ampoules, vials or straws, no embryos of a lesser health status should be processed.

3. The processing laboratory should be protected against rodents and insects.

4. The processing laboratory should be constructed with materials which permit its effective cleansing and disinfection. This should be done frequently, and following always before and after each occasion on which embryos for export are processed.

5. The laboratory must not be situated in an infected zone with regard to foot and mouth disease (except for the collection of in vivo derived bovine embryos), rinderpest, peste des petits ruminants, contagious bovine pleuropneumonia, African horse sickness, African swine fever and classical swine fever.

Article 4.7.4.

Conditions applicable to the introduction of donor animals

1. Donor animals

   a) The Veterinary Authority should have knowledge of, and authority over, the herd/flock of origin from which the donor animals have been sourced.

   b) At the time of collection, donor animals should be clinically inspected by a veterinarian responsible to the team veterinarian and certified to be free of clinical signs of diseases not included in Category 1 of the IETS classification.
Annex IX (contd)

eb) The herd of origin must not be situated in a herd / flock subject to veterinary restrictions for an infected zone for in the 30 days (60 days in the case of camelids) before and after embryo collection, with regard to foot and mouth disease (except for the collection of in vivo derived bovine embryos), rinderpest, paste des petits ruminants, contagious bovine pleuropneumonia, African horse sickness, African swine fever and classical swine fever OIE listed disease or pathogens for relevant species (see Chapter 1.2. of this Terrestrial Code), other than those that are in IETS Category 1 for the species of embryos being collected (see Article 4.7.14., and footnote).

dc) At the time of collection, the donor animals should not have been imported from another country during the previous 60 days and should have been in the herd of origin for at least 30 days prior to collection be clinically inspected by the team veterinarian, or by a veterinarian responsible to the team veterinarian and certified to be free of clinical signs of diseases other than those included in Category 1 of the IETS classification\(^1\) for the species of embryos being collected.

2. Semen donors

a) Semen used to inseminate donor animals artificially should have been produced and processed in accordance with the provisions of Chapter 4.5. or Chapter 4.6., as relevant.

b) When the donor of the semen used to inseminate donor females for embryo production is no longer living dead, and when the health status of the semen donor concerning a particular infectious disease or disease of concern was not known at the time of semen collection, additional tests may be required of the inseminated donor female after embryo collection to verify that these infectious diseases were not transmitted. An alternative may be to subject an aliquot of semen from the same collection date to testing.

c) Where natural service or fresh semen is used, donor sires should meet the same health requirements as donor females, conditions set out in Chapter 4.5, as appropriate to the species.

Article 4.7.5.

Risk management

With regard to disease transmission, transfer of in vivo derived embryos is a very low risk method for moving animal genetic material. Irrespective of animal species, there are three phases in the embryo transfer process that determine the final level of risk:

1. The first phase, which is applicable to diseases not included in Category 1 of the IETS classification categorisation\(^1\) (Article 4.7.14.), comprises the risk potential for embryo contamination and depends on:

   a) the disease situation in the exporting country and/ or zone;

   b) the health status of the herds/ flocks and the donors from which the embryos are collected;

   c) the pathogenic characteristics of the specified disease agents that are of concern to the Veterinary Authority of the importing country.

2. The second phase covers risk mitigation by use of internationally accepted procedures for processing of embryos which are set out in the IETS Manual\(^2\). These include the following:

   a) The embryos must be washed at least ten times with at least 100-fold dilutions between each wash, and a fresh pipette must be used for transferring the embryos through each wash.
b) Only embryos from the same donor should be washed together, and no more than ten embryos should be washed at any one time.

c) Sometimes, for example when inactivation or removal of certain viruses (e.g. bovine herpesvirus-1, and Aujeszky’s disease virus) is required, the standard washing procedure should be modified to include additional washes with the enzyme trypsin, as described in the IETS Manual.

d) The zona pellucida of each embryo, after washing, must be examined over its entire surface area at not less than 50X magnification to ensure that it is intact and free of adherent material.

[NOTE: All shipments of embryos must be accompanied by a statement signed by the team veterinarian certifying that these embryo processing procedures have been completed.]

3. The third phase, which is applicable to diseases not included in Category 1 of the IETS classification categorisation (Article 4.7.14.) and which are of concern to the Veterinary Authority of the importing country, encompasses the risk reductions resulting from:

a) post-collection surveillance of the donors and donor herds / flocks based on the recognized incubation periods of the diseases of concern to determine retrospectively the health status of donors whilst the embryos are stored (in species where effective storage by cryopreservation is possible) in the exporting country;

b) testing of embryo-collection (flushing) fluids and non-viable embryos, or other samples such as blood, in a laboratory for presence of specified disease agents.

Article 4.7.6.

Conditions applicable to the collection and storage of embryos

1. Media

Any biological product of animal origin used in the media and solutions for collection, processing, washing or storage of embryos should be free of pathogenic micro-organisms. Media and solutions used in the collection, freezing and storage of embryos should be sterilized by approved methods according to the IETS Manual and handled in such a manner as to ensure that sterility is maintained.

Antibiotics should be added to collection, processing, washing and storage media as recommended in the IETS Manual.

2. Equipment

a) All equipment used to collect, handle, wash, freeze and store embryos should ideally be new or at least sterilized prior to use as recommended in the IETS Manual.

b) Used equipment should not be transferred between countries for re-use by the embryo collection team only if cleaning and disinfection procedures appropriate to the disease risk concerned are followed.
Optional tests and treatments

1. The examination testing of embryos and collection or washing fluids samples can be requested by an importing country. Tests may be carried out on these samples to confirm the absence of pathogenic organisms that may be transmitted via in vivo derived embryos or to help assess whether the degree of quality control of the collection team (with regard to adherence to procedures as described in the IETS Manual) is at an acceptable level. Samples may include:

   a) Non-viable embryos/oocytes

      Where the viable, zona pellucida intact embryos from a donor are intended for export, all non-fertilized oocytes and degenerated or zona pellucida compromised embryos collected from that donor should be washed according to the IETS Manual and pooled for possible testing if requested by the importing country. Only Non-viable embryos/oocytes from one donor should be processed simultaneously and stored together.

   b) Embryo collection (flushing) fluids

      The collection fluid should be placed in a sterile, closed container and, if there is a large amount, it should be allowed to stand undisturbed for one hour. The supernatant fluid should then be removed and the bottom 10-20 ml, along with accumulated debris, decanted into a sterile bottle. If a filter is used in the collection of embryos/oocytes then any debris that is retained on the filter must be rinsed off into the retained fluid.

   c) Washing fluids

      The last four washes of the embryos/oocytes (washes 7, 8, 9 and 10) should be pooled (IETS Manual).

   d) Samples

      The samples referred to above should be stored at 4°C and tested within 24 hours. If this is not possible, then samples should be stored frozen at -70°C or lower.

2. When treatment of the viable embryos is modified to include additional washings with the enzyme trypsin (see paragraph 2c in Article 4.7.5.), the procedure should be carried out according to the IETS Manual and. Enzyme treatment is necessary only when pathogens for which the IETS recommends this additional treatment (such as with trypsin) may be present. It should be noted that such enzymatic treatment is not necessarily always beneficial and it should not be regarded as a general disinfectant. It may also have adverse effects on embryo viability, for instance in the case of equine embryos where the embryonic capsule could be damaged by the enzyme.

Conditions applicable to the storage, quarantine and transport of embryos

1. Embryos should be frozen in fresh liquid nitrogen and then stored in fresh liquid nitrogen in cleaned and disinfected tanks or containers.

2. The embryos for export should be stored in sealed sterile ampoules, vials or straws under strict hygienic conditions at a storage place approved by the Veterinary Authority of the exporting country where there is no risk of contamination of the embryos.
Annex IX (contd)

32. In case of ruminants, only embryos from the same individual donor should be stored together in the same ampoule, vial or straw.

3. The embryos should, if possible, depending on the species, be frozen, stored with fresh liquid nitrogen in cleaned and sterilized tanks or containers under strict hygienic conditions at the approved storage place.

4. Ampoules, vials or straws must be sealed at the time of freezing (or prior to export where cryopreservation is not possible), and they should be clearly identified by labels according to the standardised system recommended in the IETS Manual.

5. Liquid nitrogen containers should be sealed under the supervision of the Official Veterinarian prior to shipment from the exporting country.

6. Embryos must not be exported until the appropriate veterinary certification documents are completed.

Article 4.7.8.bis

Procedure for micromanipulation

When micromanipulation of the embryos is to be carried out, this should be done after completion of the treatments described in paragraph 2 of Article 4.7.5. and conducted in accordance with Chapter 4.9.

Article 4.7.9.

Specific conditions applicable to porcine embryos

The herd of origin should be free of clinical signs of swine vesicular disease, brucellosis and pathogenic Teschovirus encephalomyelitis.

[NOTE. The development of effective cryopreservation methods for the storage of zona pellucida-intact porcine embryos is still at a very early stage.]

Article 4.7.10.

Specific conditions applicable to ovine/caprine embryos

The herd of origin should be free of clinical signs of sheep pox, goat pox, brucellosis and bluetongue.

Article 4.7.11.

Specific conditions / comments applicable to equine embryos

The recommendations apply principally to embryos from animals continuously resident in national equine populations and therefore may be found to be unsuitable for those from equines routinely involved in events or competitions at the international level. For instance, in appropriate circumstances horses travelling with an international veterinary certificate (e.g. competition horses) may be exempt from this condition where mutually agreed upon on a bilateral basis between the respective Veterinary Authorities.
Article 4.7.12.

Specific conditions / comments applicable to camelid embryos

South American camelid embryos recovered from the uterine cavity by the conventional non-surgical flushing technique at 6.5 to 7 days post-ovulation are almost invariably at the hatched blastocyst stage, and thus the zona pellucida has already been shed. Since the embryos do not enter the uterus and cannot be recovered before 6.5 to 7 days, it would be unrealistic to stipulate for South American camelids that only zona pellucida-intact embryos can be used in international trade. It must also be noted however that, in 2008 the development of cryopreservation methods for storage of camelid embryos is still at a very early stage, and also that pathogen interaction studies with South American camelid embryos have not yet been carried out.

The herd of origin of donor animals should be free of clinical signs of foot and mouth disease, vesicular stomatitis, bluetongue, brucellosis and tuberculosis.

[NOTE: The development of cryopreservation methods for camelid embryos is still at a very early stage.]

Article 4.7.13.

Specific conditions / comments applicable to cervid embryos

The recommendations apply principally to embryos derived from animals continuously resident in national domestic or ranched cervid populations and therefore may be found to be unsuitable for those from cervids in feral or other circumstances related to biodiversity or germplasm conservation efforts.

The herd of origin should be free of clinical signs of brucellosis and tuberculosis.

Article 4.7.14.

Recommendations regarding the risk of disease transmission via in vivo derived embryos

The IETS has categorised the following diseases and pathogenic agents into four categories, which applies only to in vivo derived embryos.

1. Category 1
   a) Category 1 diseases or pathogenic agents are those for which sufficient evidence has accrued to show that the risk of transmission is negligible provided that the embryos are properly handled between collection and transfer according to the IETS Manual;
   b) The following diseases or pathogenic agents are in category 1:
      - Aujeszky's disease (pseudorabies) (swine): trypsin treatment required
      - Bluetongue (cattle)
      - Bovine spongiform encephalopathy (cattle)
      - Brucella abortus (cattle)
      - Enzootic bovine leukosis
      - Foot and mouth disease (cattle)
      - Infectious bovine rhinotracheitis: trypsin treatment required.
Annex IX (contd)

2. **Category 2**
   
a) Category 2 diseases are those for which substantial evidence has accrued to show that the risk of transmission is negligible provided that the embryos are properly handled between collection and transfer according to the IETS Manual, but for which additional transfers are required to verify existing data.

b) The following diseases are in category 2:
   - Bluetongue (sheep)
   - Caprine arthritis/encephalitis
   - Classical swine fever (hog cholera)
   - Scrapie (sheep).

3. **Category 3**
   
a) Category 3 diseases or pathogenic agents are those for which preliminary evidence indicates that the risk of transmission is negligible provided that the embryos are properly handled between collection and transfer according to the IETS Manual, but for which additional in vitro and in vivo experimental data are required to substantiate the preliminary findings.

b) The following diseases or pathogenic agents are in category 3:
   - Bovine immunodeficiency virus
   - Bovine spongiform encephalopathy (goats)
   - Bovine viral diarrhea virus (cattle)
   - Campylobacter fetus (sheep)
   - Foot and mouth disease (swine, sheep and goats)
   - Haemophilus somnus (cattle)
   - Maedi-visna (sheep)
   - Mycobacterium paratuberculosis (cattle)
   - N. ovis caninum (cattle)
   - Ovine pulmonary adenomatosis
   - Porcine reproductive and respiratory disease syndrome (PRRS)
   - Rinderpest (cattle)
   - Swine vesicular disease.

4. **Category 4**
   
a) Category 4 diseases or pathogenic agents are those for which studies have been done, or are in progress, that indicate:
   
i) that no conclusions are yet possible with regard to the level of transmission risk; or

ii) the risk of transmission via embryo transfer might not be negligible even if the embryos are properly handled according to the IETS Manual between collection and transfer.
b) The following diseases or pathogenic agents are in category 4:

- African swine fever
- Akabane (cattle)
- Bovine anaplasmosis
- Bluetongue (goats)
- Border disease (sheep)
- Bovine herpesvirus-4
- Chlamydia psittaci (cattle, sheep)
- Contagious equine metritis
- Enterovirus (cattle, swine)
- Equine rhinopneumonitis
- Escherichia coli 09:K99 (cattle)
- Leptospira borgpetersenii serovar hardjopovis (cattle)
- Leptospira sp. (swine)
- Mycobacterium bovis (cattle)
- Mycoplasma spp. (swine)
- Ovine epididymitis (Brucella ovis)
- Parainfluenza-3 virus (cattle)
- Parvovirus (swine)
- Porcine circovirus (type 2) (pigs)
- Scrapie (goats)
- Tritrichomonas foetus (cattle)
- Ureaplasma/ Mycoplasma spp. (cattle, goats)
- Vesicular stomatitis (cattle, swine).

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1 Based on available research and field information, the Research Subcommittee of the Health and Safety Advisory Committee (HASAC) of the International Embryo Transfer Society (IETS) has categorised some diseases based on their relative risk of dissemination by properly processed and handled in vivo derived embryos. This Chapter that contains the complete list of IETS categorised diseases is shown in Article 4.7.14.

CHAPTER 4.8.

COLLECTION AND PROCESSING OF IN VITRO FERTILISED BOVINE PRODUCED EMBRYOS / IN VITRO MATURING BOVINE OOCYTES FROM LIVESTOCK AND HORSES

Article 4.8.1.

Aims of control

Production of embryos in vitro involves the collection of oocytes from the ovaries of donors, in vitro maturation and fertilization of the oocytes, then in vitro culture to the morula/blastocyst stage at which they are ready for transfer into recipients. The purpose of official sanitary control of in vitro fertilised bovine produced embryos intended for movement internationally is to ensure that specific pathogenic organisms, which could be associated with such embryos, are controlled and transmission of infection to recipient animals and progeny is avoided. The conditions outlined in this Chapter are also applicable where the movement of in vitro maturing (IVM) oocytes is intended.

Article 4.8.1.bis

Conditions applicable to the embryo collection production team

The embryo production team is a group of competent technicians, including at least one veterinarian, to perform the collection and processing of bovine ovaries/oocytes and the production and storage of in vitro fertilised bovine embryos. The following conditions should apply:

1. The team should be supervised by a team veterinarian.

2. The team veterinarian is responsible for all team operations which include the hygienic collection of ovaries and oocytes and all other procedures involved in the production of embryos intended for international movement.

3. The team veterinarian should be specifically approved for this purpose by an official Veterinarian.

4. Team personnel should be adequately trained in the techniques and principles of disease control. High standards of hygiene should be practised to preclude the introduction of infection.

5. The production team must have adequate facilities and equipment for:
   a) collecting ovaries and/or oocytes;
   b) processing of oocytes and production of embryos at a permanent site or mobile laboratory;
   c) storing oocytes and/or embryos.

   These facilities need not necessarily be at the same location.

6. The collection embryo production team must keep a record of its activities, which must be maintained for inspection by the approving Veterinary Authority for a period of at least 2 years after the embryos have been exported.
7. The embryo production team should be subjected to regular inspection at least once a year by an Official Veterinarian to ensure compliance with procedures for the sanitary collection and processing of oocytes, and the production and storage of embryos.

8. The production team must not operate in an infected zone for foot and mouth disease, contagious bovine pleuropneumonia, bluetongue or rinderpest.

Article 4.8.2.

Conditions applicable to the processing laboratories

The A processing laboratory is a premises used by the embryo production team may be mobile or permanent. It may be connected to the oocyte recovery area or at a separate location. It is a facility in which oocytes which have been recovered from ovaries are then matured and fertilised, and where the resulting embryos are further cultured in vitro. It may be contiguous with the oocyte recovery area or may be at a separate location.

The embryos so produced may also be subjected to any required treatments such as washing before freezing, and storage and quarantine in this laboratory.

Additionally:

1. The laboratory should be under the direct supervision of the team veterinarian and regularly inspected by an Official Veterinarian.

2. While embryos for export are being produced prior to their storage in ampoules, vials or straws, no oocyte/embryo of a lesser health status should be recovered or processed in the same laboratory.

3. The laboratory should be protected against rodents and insects.

4. The processing laboratory should be constructed with materials which permit its effective cleaning and disinfection. This should be done frequently and always before and after each occasion on which embryos for export are processed.

5. The laboratory must not be situated in an infected zone for foot and mouth disease, contagious bovine pleuropneumonia, bluetongue or rinderpest.

Article 4.8.3.

Conditions applicable to the introduction of donor animals

Oocytes for the in vitro production of IVF embryos are obtained from donors basically in one of two different ways: individual collection or batch collection. The recommended sanitary conditions for these differ.
Individual collection usually involves the aspiration of oocytes from the ovaries of individual live animals on the farm where the donor animal resides, or at the laboratory. Occasionally oocytes may also be recovered from individual live donors by aspiration from surgically excised ovaries. When oocytes are recovered from individual live animals, the procedures for these donors should follow the recommendations set out in Article 4.7.4. When oocytes are recovered from individual live animals, the conditions for these donors should resemble those set out in Article 4.7.4.

In these cases, cleaning and sterilisation of equipment (e.g., ultrasound guided probes) is especially important and must be carried out between each donor in accordance with the recommendations in the Procedures Manual of the International Embryo Transfer Society (IETS).

Batch collection usually involves the removal of ovaries from batches of donor slaughtered animals at an abattoir; but may alternatively involve the surgical removal of ovaries from live donors, these ovaries are then transported to the processing laboratory where the oocytes are removed from the ovarian follicles by aspiration. Batch collection involving abattoir derived ovaries has the disadvantage that it is usually impractical to relate the ovaries which are transported to the laboratory to the donors which were slaughtered at the abattoir. Nevertheless, it is critical to ensure that only healthy tissues are obtained and that they are removed from the donors and transported to the laboratory in a hygienic manner.

Additionally:

1. The Veterinary Authority should have knowledge of, and authority over, the herd(s) / flock(s) of origin of from which the donor animals have been sourced.

2. The donor females animals should not originate from herds / flocks which are subject to veterinary restrictions for listed diseases — foot and mouth disease, contagious bovine pleuropneumonia, bluetongue or rinderpest — and neither should the removal of any tissue or aspiration of oocytes should not take place in an infected zone or one that is subject to veterinary restrictions for listed disease — foot and mouth disease, contagious bovine pleuropneumonia, bluetongue or rinderpest.

3. In the case of oocyte recovery from individual animals live donors, post-collection surveillance of the donors and donor herds / flocks should be conducted based on the recognized incubation periods of the diseases of concern to determine retrospectively the health status of donors should be conducted.

4. In the case of oocyte recovery from batches of ovaries collected from an abattoir, The abattoir should be officially approved and under the supervision of a veterinarian whose responsibility it is to ensure that ante-mortem and post-mortem inspections of potential donor animals are carried out, and to certify them to be free of clinical or pathological signs of contagious disease of concern transmissible to cattle that may be transmissible by bovine semen or embryos.

5. The donor females animals slaughtered at an abattoir should not have been designated for compulsory slaughter for a notifiable disease and other animals of a lesser health status must should not be slaughtered at the same time as donors from which ovaries and other tissues will be removed.

6. Records of the identities and origins of all donors must be kept should be maintained for inspection by the approving authority for a period of at least 2 years after the embryos have been exported.
6. Batches of ovaries and other tissues collected from an abattoir should not be transported to the processing laboratory before confirmation has been obtained that ante- and post-mortem inspection of donors has been satisfactorily completed.

7. Equipment for the removal and transport of ovaries and other tissues should be cleaned and sterilized before use and exclusively used for these purposes.

8. Records of the identities and origins of all donors should be maintained for inspection by the Veterinary Authority for a period of at least 2 years after the embryos have been exported. While this may be difficult to achieve in the case of batch collection, it is to be expected that the identities of the herds/flocks from which the donors originated will be maintained.

Article 4.8.4.

Testing of oocytes, embryos, semen and culture media Optional tests and treatments

The main approach for ensuring that IVF in vitro produced embryos do not transmit disease is by testing various materials to confirm the absence of pathogenic organisms which are of concern to the importing country are free of pathogenic organisms is the testing of non-viable oocytes/embryos and associated co-culture cells, fluid and media.

Tests may also be used to assess whether quality control procedures being applied in the processing laboratory are of an acceptable standard.

Tests may be carried out on the following materials to confirm the absence of pathogenic organisms that may be transmissible by bovine semen or embryos and that are of concern to the importing country:

a) non-viable oocytes/embryos: all non-viable oocytes/embryos at any stage of the in vitro production line from batches intended for export should be pooled for testing.

b) samples of in vitro maturation medium taken prior to mixing the oocytes with semen for the fertilisation process;

c) samples of embryo culture medium taken immediately prior to embryo storage.

These samples should be stored at 4°C and tested within 24 hours. If this is not possible, then the samples should be stored frozen at -70°C or lower.

In the case of oocyte recovery from individual animals or batch collection from live donors, monitoring of clinical health status and post collection testing of donors for diseases of concern may be considered.

Additionally:

1. Semen used to fertilise oocytes in vitro should meet the health requirements and standards set out in Chapter 4.5 as appropriate to the species.

When the donor of the semen used to fertilise the oocytes is no longer living, and when the health status of the semen donor concerning a particular infectious disease or diseases of concern was not known at the time of semen collection, additional tests on the spare IVF embryos may be required to verify that these infectious diseases were not transmitted. An alternative may be to subject test an aliquot of semen from the same collection date to testing.
2. Any biological product of animal origin, including co-culture cells and media constituents, used in oocyte recovery, maturation, fertilisation, culture, washing and storage should be free of living pathogenic micro-organisms. Media should be sterilised prior to use by approved methods according to the IETS Manual and handled in such a manner as to ensure that sterility is maintained. Antibiotics should be added to all fluids and media as recommended in the IETS Manual.

3. All equipment used to recover, handle, culture, wash, freeze and store oocytes/embryos should be new or cleaned and sterilised prior to use as recommended in the IETS Manual.

**Article 4.8.4.bis**

**Risk management**

With regard to disease transmission, transfer of in vitro produced embryos is a low risk method for moving animal genetic material although the risk is not quite as low as for in vivo derived embryos. It should be noted that categorisation of diseases/disease agents by the IETS, as described for in vivo derived embryos in Article 4.7.14, does not apply in the case of in vitro produced embryos. Irrespective of the animal species, there are three phases in the embryo production and transfer process that determine the final level of risk. These are as follows:

1. The first phase comprises the risk potential for ovary/oocyte/embryo contamination and depends on:
   a) the disease situation in the exporting country and/or zone;
   b) the health status of the herds/flocks and the donors from which the ovaries/oocytes/embryos are collected;
   c) the pathogenic characteristics of the specified disease agents that are of concern to the Veterinary Authority of the importing country;

2. The second phase covers risk mitigation by the use of internationally accepted procedures for the processing of embryos which are set out in the IETS Manual. These include the following:
   a) after the in vitro culture period is finished the embryos should be washed at least ten times with at least 100-fold dilutions between each wash, and a fresh pipette should be used for transferring the embryos through each wash;
   b) only embryos from the same donor (in the case of individual collection) or from the same batch (in the case of batch collection) should be washed together, and no more than ten embryos should be washed at any one time;
   c) sometimes, for example when inactivation or removal of certain viruses (e.g. bovine herpesvirus-1, or Aujeszky’s disease virus) is required, the standard washing procedure should be modified to include additional washes with the enzyme trypsin, as described in the IETS Manual;
   d) the zona pellucida of each embryo, after washing, should be examined over its entire surface area at not less than 50X magnification to ensure that it is intact and free of adherent material;

3. The third phase, which is applicable to diseases which are of concern to the Veterinary Authority of the importing country, encompasses the risk reductions resulting from:
Annex IX (contd)

a) post-collection surveillance of the donors and donor herds/flocks based on the recognised incubation periods of the diseases of concern to determine retrospectively the health status of the donors whilst the embryos are stored (in species where effective storage by cryopreservation is possible) in the exporting country. Post-collection surveillance of donors is not, of course, possible in the case of batch collection from an abattoir, although surveillance of the herds/flocks of origin may be possible.

b) testing of oocytes/embryos, co-culture cells, media and other samples (e.g. blood) (as referred to in Article 4.8.4.) in a laboratory for presence of disease agents.

Article 4.8.5.

Conditions applicable to the processing, storage, quarantine and transport of embryos/ova

1. After the culture period is finished but prior to freezing, storage and transport, the embryos should be subjected to washing and other treatments similar to those specified for in vivo derived embryos in accordance with the IETS Manual.

2. Only embryos from the same individual donor, in the case of individual animal recovery, or from the same batch collection, should be washed together, stored together in the same ampoule, vial or straw.

3. The zona pellucida of each embryo must be examined over its entire surface area at not less than 50X magnification and certified to be intact.

4. The IVF embryos should if possible, depending on the species, be stored in sealed sterile ampoules, vials or straws and then frozen in fresh liquid nitrogen or other cryoprotectant and then stored in fresh cryoprotectant in cleaned and sterilised tanks or containers under strict hygienic conditions at a storage place, approved by the Veterinary Authority of the exporting country, where no risk of contamination of the embryos can occur.

5. Only embryos from the same individual donor or batch collection should be stored together in the same ampoule, vial or straw.

6. Ampoules, vials or straws must be sealed at the time of freezing and should be labelled according to the IETS Manual.

7. Liquid nitrogen containers should be sealed prior to shipment from the exporting country.

8. Embryos must not be exported until the appropriate veterinary certification documents are completed.
Article 4.8.6.

Procedure for micromanipulation

When micromanipulation of the embryos is to be carried out, this should be done after completion of the treatments described in Article 4.8.5., 4bis paragraph 2, and conducted in accordance with Chapter 4.79.

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1. Where transportation of in vitro maturing (IVM) oocytes is intended, the conditions outlined in this Chapter are also applicable.


   An up-to-date list of relevant scientific publications is available on the IETS Website Homepage at http://www.iets.uiue.edu where the link entitled ‘Embryo-Pathogen Research and Reference Lists’ may be visited.

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CHAPTER 4.9.

COLLECTION AND PROCESSING OF MICROMANIPULATED BOVINE EMBRYOS / OOCYTES FROM LIVESTOCK AND HORSES

Article 4.9.1.

Introduction

Neither Chapter 4.7, which recommends official sanitary control measures for the international movement of intact, in vivo derived bovine embryos, nor likewise Chapter 4.8, which recommends measures for in vitro fertilized produced bovine embryos in vitro maturing bovine oocytes. Neither of those Chapters covers embryos which have been subjected to biopsy, splitting, transgene injection, intracytoplasmic sperm injection (ICSI), nuclear transplantation transfer or other micromanipulations interventions which breach the integrity of the zona pellucida. Such embryos oocytes are subsequently those referred to here as having been 'micromanipulated embryos'.

It should be noted that complete removal of granulosa cells prior to micromanipulation of oocytes, zygotes and embryos or other adherent material from the outer surface of the zona pellucida of oocytes, zygotes and embryos is necessary prior to micromanipulation to avoid lowering their health status.

Removal of such material from the zona pellucida of immature oocytes can be difficult. However, to bring micromanipulated embryos oocytes within the scope of the above mentioned Chapters, the following conditions shall apply:

Article 4.9.2.

1. Prior to any micromanipulation which involves breaching the zona pellucida, all embryos oocytes must should be collected and processed according to the sanitary conditions laid down in Chapter 4.7. (in vivo derived embryos) or produced according to the sanitary conditions laid down in Chapter 4.8. (in vitro fertilised produced bovine embryos in vivo maturing bovine oocytes).

2. Responsibility for the embryos oocytes must remains with the embryo collection team (in vivo derived embryos) or with the embryo production team (in vitro fertilised bovine produced embryos), and all processing involving micromanipulation should be carried out in an approved processing laboratory under supervision of an approved team veterinarian (see Articles 4.7.2. and 4.7.3., and Articles 4.8.1. and 4.8.2., as relevant appropriate).

3. Donor animals must should comply with the conditions laid down in Article 4.7.4. (in vivo derived embryos) or Article 4.8.3. (in vitro produced fertilised bovine embryos in vivo maturing bovine oocytes), whichever is appropriate. The Risk management and criteria for testing samples to ensure that embryos are free of pathogenic organisms are laid down in Articles 4.7.5. and 4.7.7. and in Articles 4.8.4. and 4.8.5., respectively, and these should be followed.
4. All embryos to be micromanipulated must be washed according to the protocols laid down in the IETS Manual (1998) and they must be observed to have an intact zona pellucida before and after washing. Only embryos from the same donor, or, in the case of some in vitro produced embryos (see Chapter 4.8.) embryos originating from the same batch, collection of ovaries from an abattoir (see Chapter 4.8.), should be washed together at the same time. After washing, but before micromanipulation, the zona pellucida of each embryo should be examined over its entire surface area at not less than 50X magnification and certified to be intact and free of adherent material.

5. If surrogate zonae are used, they should be of bovine origin from the same species and the embryos/oocytes from which they are obtained should be treated in the same manner as if they were in vivo derived or in vitro produced embryos intended for international movement.

**Article 4.9.3.**

**Procedures for micromanipulation**

The term ‘micromanipulation’ covers several different procedures and a variety of specialized microsurgical instruments and other equipment may be used. However, from the standpoint of animal health, any cutting, penetrating or breaching of the integrity of the zona pellucida is an action that can alter the health status of an embryo. To maintain health status during and after micromanipulation, the following conditions should apply:

1. **Media**

   Any product of animal origin, including co-culture cells and media constituents, used in the collection or production of embryos, oocytes or other cells, and in their micromanipulation, culture, washing and storage should be free of pathogenic micro-organisms (including transmissible spongiform encephalopathy agents, sometimes called prions). All media and solutions should be sterilized by approved methods according to the IETS Manual and handled in such a manner as to ensure that sterility is maintained. Antibiotics should be added to all fluids and media as recommended in the IETS Manual.

2. **Equipment**

   Equipment (e.g. microsurgical instruments which have direct contact with embryos) should either be of the single-use type (disposed of after each embryo / oocytes batch) or should be effectively sterilised between embryos / oocytes batch in accordance with recommendations in the IETS Manual.

3. **Nuclei for transfer transplantation (‘nuclear transfer’)**

   a) Where it is intended to transplant nuclei derived from pre-hatching stage (i.e. zona pellucida intact) embryos, the parent embryos from which those nuclei are derived should fulfil the conditions of this Chapter. Where nuclei derived from other types of donor cell (e.g. post-hatching stage embryos, embryonic, fetal and adult cells, including spermatozoa/spermatids for ICSI) are to be transplanted, the parent embryo, fetus or animal from which those donor cells originate, and the methods whereby they are derived, including cell culture, should comply with the relevant animal health standards recommended elsewhere in this Terrestrial Code and in the Terrestrial Manual.

   b) Where it is intended to transplant a nucleus into an intact oocyte (e.g. for ICSI), or into an enucleated oocyte (for nuclear transfer), those oocytes should be collected, cultured and manipulated according to the recommendations in this Chapter and/or in Chapter 4.7.

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Optional tests and treatments

The importing country may request that tests be carried out on certain samples or that embryos are treated to ensure that specified pathogenic organisms are absent.

1. **Samples**

   Samples to be tested may include those referred to in Article 4.7.7. and/or in Article 4.8.4. Where cells other than from zona pellucida-intact embryos (e.g. somatic or sperm cells) are used as donors of nuclei for transplantation, then samples or cultures of those donor cells may also be tested.

2. **Treatments**

   Treatments of embryos with the enzyme trypsin or other substances proven to inactivate or remove pathogenic organisms, and which are harmless to the embryo, may be requested when pathogens that are not removed by routine washing may be present, but also be applied prior to any micromanipulation, and according to the IETS Manual.

Article 4.9.5.

Conditions applicable to storage, quarantine and transport

Micromanipulated embryos should be stored, quarantined and transported according to the conditions laid down in Article 4.7.8. or in points 4, 5, 6, 7 and 8 of Article 4.8.5. Veterinary certification documents should identify all micromanipulations, where and when they were carried out.

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2 If the samples mentioned above in point 1. of Article 4.9.4. are to be tested for pathogenic agents, then the microbiological techniques in current use for those agents would be appropriate.
CHAPTER 4.10.

COLLECTION AND PROCESSING OF LABORATORY RODENT AND RABBIT EMBRYOS / OVA

Article 4.10.1.

Conditions applicable to the maintenance of laboratory animal colonies

Maintenance of laboratory animal colonies of specific genotypes requires intensive breeding management within specialised premises. They may be kept in a gnotobiotic environment, in either a ‘germfree’ system or a ‘barrier’ room (usually with defined flora), in a conventional colony, or under undefined conditions. In both the germfree and barrier systems, the animals are raised in a controlled environment according to protocols that attempt to eliminate potential sources of microbiological contamination. The primary difference is that the barrier maintained animals have been inoculated with known (defined) microbes using a cocktail of non-pathogenic flora, whereas germfree animals are kept free from both pathogenic and non-pathogenic microbes.

A second category is where laboratory animals are kept in closed, conventional colonies within which known pathogens may exist. Here, less rigid colony management protocols are used to control potential sources of contamination, but implementation of simple aseptic precautions (e.g. autoclaving of food and bedding) should allow animals to be maintained in a microbiologically defined system. Finally, laboratory animals may live in environments with undefined microbiological conditions (e.g. non-restricted colonies, free ranging animals).

Disease testing and donor animal/embryo handling requirements can therefore be considered as being of three distinct types, depending on the type of colony being dealt with, i.e. defined floral, conventional and undefined. The health status of all colonies should be confirmed quarterly by bacteriological, virological, parasitological, serological and immunohistochemical tests on pre-designated sentinel animals or other representative animals of the colony (e.g. older breeding males which have sired multiple litters).

Microbial status of laboratory animal colonies

Colonies of the various species and genotypes of laboratory animals are usually kept within specialised premises and their microbial status depends largely on the system whereby the colony was formed and is maintained. In this Chapter the microbial status of colonies is considered to be of three main types: defined, conventional and undefined. Colonies of defined status are those where, at least initially, the animals are totally free of pathogenic and non-pathogenic micro-organisms (i.e. gnotobiotic), although sometimes a cocktail of known, non-pathogenic micro-organisms has been given subsequently. In either case defined colonies are kept in highly controlled environments in barrier maintained rooms, with strict protocols in place to exclude all potential sources of unwanted microbiological contamination. Colonies of conventional status are those where the animals are kept in closed colonies but where known (specific) pathogens as well as non-pathogenic micro-organisms may exist. While management protocols for conventional colonies may be less rigid than those for defined colonies, they are designed to control potential sources of microbial contamination. Simple aseptic precautions (e.g. the autoclaving of food and bedding) are taken to ensure that the animals do not become infected with any unwanted microflora. Finally, laboratory animals may be kept in microbiologically undefined colonies which are unrestricted and may include free ranging animals. Details of these different types of colony can be found in the FELASA Report.

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The health status of defined and conventional colonies should be confirmed at least quarterly by bacteriological, virological, parasitological, serological and other tests on pre-designated sentinel animals or other representative members of the colony. Older breeding males which have sired multiple litters are often selected for this purpose.

The purpose of official sanitary control of laboratory rodent and rabbit embryos intended for movement internationally is to ensure that specific pathogenic micro-organisms, which could be associated with such embryos, are controlled and transmission of infection to recipient animals, progeny and colonies, is avoided. Requirements for the management of donors and processing of embryos vary depending on the microbial status of the colony, i.e. whether it is defined (including gnotobiotic), conventional, or undefined.

Article 4.10.2.

Conditions applicable to the embryo production team/laboratory

The embryo collection team is a group of competent technicians including at least one experienced professional to perform the collection, processing and storage of embryos/oocytes.

The following conditions should apply:

1. The embryo production team must be composed of competent technicians supervised by an experienced embryologist holding a graduate academic degree (e.g. M.S., Ph.D., D.V.M.).

2. The team professional is responsible for all team operations which include verification of colony and donor health status, sanitary handling and surgery of donors, disinfection and hygienic procedures. The team professional should be responsible to the institute veterinarian.

3. The institute veterinarian should be certified or accredited in laboratory animal care and should be specifically approved for the purpose of embryo collection for export. It is the responsibility of the institute veterinarian to ensure that required health profiling procedures appropriate for the colony status are implemented. He/she is responsible for certifying that the embryo handling procedures and laboratory facilities conform to the requirements laid down in this Chapter.

4. Team personnel should be adequately trained in the techniques and principles of disease control and in the use of aseptic techniques in embryo handling. Laboratory sanitary procedures must conform with requirements in the IETS Manual. The zoonotic potential of specific pathogens affecting the various laboratory animal species should be identified and understood so as to avoid contamination of colonies via human vectors, and vice versa.

5. The embryo production team must use all necessary precautions to protect the animals, animal facilities, laboratory and equipment against microbiological contamination. In particular, the zoonotic potential of specific pathogens should be identified and understood by staff members to avoid contamination of colonies via human vectors, or vice versa. High standards of hygiene should be practiced to preclude the introduction of infection to the donor animals, colonies, facilities, and equipment. Restrictions should be established to prevent free access of personnel into the embryo collection and handling laboratory facilities especially after their exposure such personnel have been exposed to other animal facilities.
6. The team should have adequate facilities and equipment for:
   a) collecting embryos;
   b) processing and treatment of embryos at a permanent site or mobile laboratory;
   c) storing embryos.

4. Proper records must be maintained for inspection by the chief embryologist (i.e. supervisor).

Until standardised record sheets are developed for laboratory animals, it is the responsibility of each laboratory to maintain complete animal and embryo records (i.e. embryo collection, cryopreservation data). Information of the type shown in standard IETS record sheets for livestock species should be incorporated, where applicable, and data such as embryo quality grading system, morphological stage at cryopreservation and genotypic identification of the donors should be clearly given in the records.

5. It is the responsibility of the chief embryologist (i.e. laboratory supervisor) to ensure that the complete animal and embryo records, including records of collection, processing and storage of embryos are maintained. In addition, the containers must be correctly identified using a standard format which includes embryo species/genotype, cryopreservation date, number and stage of embryos, container number and indication of any specialised procedure (e.g. in vitro fertilisation, micromanipulation) or condition (e.g. germfree, microbiologically defined). Record sheets of the type shown in the IETS Manual for livestock species should be used where applicable, and data such as genotypic identification of the donors, embryo quality grading, morphological stage and condition should be given. If appropriate the embryo collection team should keep a record of its activities which should be maintained for inspection by the Veterinary Authority for at least 2 years after the embryos have been exported.

8. The embryo collection team, if involved in the export of embryos, should be subject to regular inspection, preferably annually, by an Official Veterinarian to ensure compliance with procedures for the sanitary collection, processing and storage of embryos.

**Article 4.10.2bis**

**Conditions applicable to the processing laboratory**

A processing laboratory used by the embryo collection team is a facility in which embryos are recovered from donors (or from their excised reproductive tracts), and from the collection media. Here also the embryos are examined and subjected to any required treatments such as washing, cryopreservation for storage and quarantine pending results of any diagnostic procedures. The processing laboratory may be part of a specifically designed collection and processing unit, or a suitably adapted part of an existing building. It may be on the premises where the donor animals are kept but in this case should be physically separated from animals.

Additionally:

1. The processing laboratory should be under the supervision of the institute veterinarian and be inspected by an Official Veterinarian.
Annex IX (contd)

2. While embryos for export are being handled prior to their storage in ampoules, vials or straws, no embryos of lesser health status should be processed.

3. The processing laboratory should be constructed with materials which permit its effective cleansing and disinfection. This should be done frequently, and always before and after each occasion on which embryos for export are processed.

Article 4.10.2tris

Risk management

With regard to disease transmission, transfer of in vitro derived embryos is a very low risk method for moving the genetic material of laboratory animals. Irrespective of animal species, there are three phases in the embryo transfer process that determine the final level of risk:

1. The first phase comprises the risk potential for embryo contamination and depends on:
   a) the disease situation in the exporting country and/or zone;
   b) the microbial status of the colony (i.e. defined, conventional or undefined) and the donors from which the embryos are collected;
   c) the pathogenic characteristics of the specified disease agents that are of concern to the Veterinary Authority of the importing country.

2. The second phase covers risk mitigation by use of internationally accepted procedures for processing of embryos which are set out in the IETS Manual. These include the following:
   a) Depending on microbial status of the colony, the embryos should be washed up to ten times with at least 100-fold dilutions between each wash, with a fresh pipette being used for transferring the embryos through each wash.
   b) Only embryos from the same donor should be washed together, and no more than ten embryos should be washed at any one time.
   c) Sometimes, for example when removal of certain viruses (e.g. herpesviruses) is required, the standard washing procedure should be modified to include additional washes with the enzyme trypsin, as described in the IETS Manual.
   d) The zona pellucida of each embryo, after washing, should be examined over its entire surface area at not less than 50X magnification to ensure that it is intact and (apart from the mucin layer in the case of rabbit embryos) free of adherent material.

3. The third phase, which is applicable to diseases of concern to the Veterinary Authority of the importing country, encompasses risk mitigation resulting from:
   a) post-collection surveillance of the microbial status of the donor colony based on the recognized incubation periods of the disease of concern to determine retrospectively the health status of the colony whilst the embryos are stored (in species where effective storage by cryopreservation is possible) in the exporting country;
   b) post-mortem testing of the donor(s) or other samples such as blood, embryo-collection (flushing) fluids and non-viable embryos, in a laboratory for presence of specified disease agents.
Article 4.10.3.

Conditions applicable to the embryo team/ institute veterinarian

1. The veterinarian, certified in laboratory animal care or laboratory animal accredited, must ensure that the required colony health profiling procedures are implemented, and the results are reviewed and properly recorded before shipment of embryos. He/ she is also responsible for confirming that proper animal management/ sanitation conditions have been maintained. It is the responsibility of the institute veterinarian to ensure that required health testing procedures are implemented to demonstrate micro-bial status of the colony (i.e. defined, conventional or undefined). Colony microbial status should be reviewed by the institute veterinarian before shipment of the embryos.

2. The veterinarian is responsible for certifying that the embryo handling procedures and laboratory conditions were maintained in accordance with the IETS Manual Articles 4.10.2. and 4.10.2bis.

3. The veterinarian must supervise all quarantine practices to protect against unwanted contamination and spread of disease, and to ensure that valid results are generated is responsible for the risk management procedures outlined in Article 4.10.2tris.

4. The veterinarian must should authorise all embryo shipments, ensuring that the correct embryo collection records and veterinary certification documents and embryo collection records are have been completed and are included in the shipments.

Article 4.10.4.

Test programmes for donor animals

Sentinel animals in each donor colony should be subjected to routine monthly microbial screening. Testing for specific pathogens is species dependent and will undoubtedly also be influenced by geographic location. Recommendations regarding specific microbial agents to be tested for in mice, rats, cotton rats, hamsters, guinea pigs, gerbils and rabbits have been published elsewhere.

Article 4.10.5.

Conditions applicable to the embryo/animal handling donors from animal colonies of different microbial status

It should be noted that the conditions applicable to donor animals vary according to the microbial status of the colony from which they originate, i.e. defined, conventional or undefined.

Sentinel animals in each donor colony of defined and conventional status should be subjected to routine microbial screening, preferably monthly, but at least quarterly. Testing for specific pathogens depends on the animal species and may be influenced by geographical location. Recommendations regarding specific microbial agents to be tested for in different laboratory animal species have been published elsewhere.

1. Defined microbial conditions status

   a) Germfree and Microbiologically defined colonies (Article 4.10.1), barrier maintained animals represent the cleanest sources of gametes, and the embryos recovered from these animals can be regarded as pathogen free.
b) Since the animals themselves male and female donors are pathogen free or possess defined flora (usually based on random, monthly testing of sentinel animals), dissection of the female reproductive tract and embryo isolation collection procedures can be performed under aseptic laboratory conditions, and do not require the use of a biological safety cabinet if appropriate.

c) Strict aseptic procedures should nevertheless be followed and, while embryo washing is not essential to safeguard against any possible air-borne contamination in the laboratory, it is recommended that embryos undergo at least a 3 step washing procedure. In each wash, embryos should be gently agitated in the medium, and the wash volume must constitute at least a one hundred-fold dilution of the volume in which the embryos are transferred. Embryo washed as described in point 2 of Article 4.10.2tris is not necessary but it is recommended that embryos are washed 2 or 3 times. In each wash, embryos should be gently agitated in the medium.

d) Microbial testing of flush or washing media is not required.

d) Cryopreserved embryos should be designated, in the appropriate records, The embryos should be recorded as coming from a germfree or microbiologically defined, barrier maintained colony, thus indicating that additional safeguards special risk management procedures (Article 4.10.2tris) for pathogen removal are not necessary. Isolation and health status monitoring of The need to quarantine the embryo recipients should be considered but the need to quarantine them is a matter for the importing laboratory institute.

2. Conventional conditions

a) Animals maintained under these conditions generally represent closed colonies whose Colonies of conventional microbial status are usually closed and their health status is routinely profiled monitored (Article 4.10.1). The animals may have been exposed to various pathogens, resulting in the isolation of infectious agents with positive antibody titres or even active clinical disease. However, prior to embryo collection there should be familiarity with the pathogen(s) of particular concern in each individual colony should be well known.

b) Reproductive tracts (uteri, oviducts and/or ovaries) should be removed at a separate site and then taken into the embryo processing laboratory. These procedures should be performed by separate different technicians or, at the very minimum, their protective clothing should be changed between locations. If the animals are to should be handled in the laboratory, the tracts should be dissected out within a biological safety cabinet. This will help protect against the possible shedding of pathogens into the laboratory itself.

c) Once the reproductive tracts have been removed, embryo recovery should be performed under aseptic conditions. Embryos must be inspected (>100x) for the presence of cracks in the zona pellucida and only zona-intact embryos should be kept. They must then be washed using the standard 10 step procedure, described. Depending on which, if any, pathogens are known to occur in the colony, embryos should be processed according to the risk management procedures, including washing, as described in Article 4.10.2tris, and in the IETS Manual. This recommendation could be waived in the future if sufficient research evidence from embryo-pathogen interaction studies warranted it.
Annex IX (contd)

d) Embryos derived from animals that have positive antibody titres or other evidence of specific pathogens should only be transferred into a new colony via a quarantine system, using microbiologically defined recipient females. As an additional safeguard, Quarantine may also be appropriate if there is any uncertainty about the donor or disease status of the embryos, quarantining of recipients should be applied the microbial status of the donor colony or the donors. In certain situations where the embryos might have been exposed to bacterial infection (e.g. mycoplasma), they should be cultured in a medium containing an appropriate antibiotic for 24 h pre-freezing or post-thawing and prior to transfer before cryopreservation, or in the interval between thawing and transfer into recipients.

e) If the embryos were not handled in the recommended manner, this must be indicated on the shipment records, and mandatory quarantining of the recipient dam and offspring should be imposed by the recipient institution until their health status is confirmed. The recipient dam should then be tested post-weaning for pathogens, and introduction of the progeny into the colony should only take place if test results are satisfactory. If the recipient institution does decide to quarantine the recipient dam and offspring until their health status is confirmed, the recipients should be tested post-weaning for pathogens of concern, and introduction of offspring into the colony should only take place if the test results are satisfactory.

3. Undefined microbial conditions

a) These animals are derived from either the wild Embryos from free ranging animals or from colonies of unknown health status and embryos from them require maximum precautions the full range of risk management procedures that are described in Article 4.10.2tris and in the IETS Manual. The health status of breeder males and donor females should be determined. The procedures resemble those used for embryos of livestock as recommended in Chapter 4.7. and Chapter 4.8. of this Terrestrial Code. Ideally, the breeder males and donor females should be separated from other animals and tested 15 days before and on the day of breeding (for males) or at embryo collection (for females). Alternatively, the animals could be incorporated into a conventional colony, where, over time, a health history can be documented to reduce the strict monitoring and embryo handling requirements.

b) A Biological safety cabinet should be used for all animal, tissue and embryo handling donors and reproductive tissues, and for processing embryos.

c) Post-mortem testing of the donor females for diseases or pathogens of concern to the importing country may be appropriate after the embryos/oocytes have been collected. Alternatively if embryos are collected surgically an aliquot of flush fluid from each donor, or a pooled sample, should be tested for the presence of specific pathogens of concern to the importing country and laboratory.

d) Embryos must be washed at least 10 times in accordance with the protocols in the IETS Manual (i.e. the 10 step wash, possibly including trypsin treatment in the case of certain herpesviruses) and an aliquot of media from the last four (pooled) washes should be tested for pathogens trypsin treatment should be used if presence of certain pathogenic herpesviruses is of concern.

e) Cryopreserved embryos must be stored in the exporting laboratory until such time as the necessary disease screening of colonies, tissues and fluids is completed and the certification documents completed and signed by the institute veterinarian.
Annex IX (contd)

9. On arrival in the importing country the All embryos from these animals must should be transferred into a colony via recipients in a quarantine system, as discussed above. Recipients should be tested at intervals appropriate to recognized incubation periods of the disease of concern. In addition to testing the recipients dam after transfer, all the offspring should be tested at 12 weeks of age and/or individuals from successive generations should be tested before their introduction into breeding colonies outside the quarantine facility.

Article 4.10.5 bis.

Conditions applicable to the storage and transport of embryos

1. Embryos for export should be frozen in fresh liquid nitrogen and then stored in fresh liquid nitrogen in cleaned and disinfected tanks or containers.

2. The embryos should be stored in sealed sterile ampoules, vials or straws under strict hygienic conditions at a storage place approved by the Veterinary Authority of the exporting country. Only embryos from the same donor should be stored together in the same ampoule, vial or straw.

3. Ampoules, vials or straws should be sealed at the time of freezing (or prior to export where cryopreservation is not possible) and they should be clearly identified according to or similar to the system recommended in the IETS Manual. Identification should include details of the species/genotype of the donors, microbial status (e.g. defined, conventional or undefined), collection/cryopreservation date, number and developmental stage of the embryos, container number and details of any specialized procedure such as in vitro fertilization, micromanipulation.

4. Liquid nitrogen storage containers should be sealed under the supervision of the Official Veterinarian prior to shipment from the exporting country.

5. Embryos should not be exported until the appropriate veterinary certificates are completed.

Article 4.10.6.

Special experimental circumstances

Procedures for in vitro fertilization and micromanipulation

If embryos are to be cryopreserved following specialised produced by in vitro fertilization of oocytes, it is advised that the washed sperm should be used so as to minimize the risk of possible pathogen exposure. If embryos are to undergo micromanipulation procedures that involve penetration of the zona pellucida, they must undergo the required washing steps (depending on colony status) before treatment. In the case of in vitro fertilisation, to minimise possible pathogen exposure, it is also advised that only washed sperm should be used. Embryos should be washed again before cryopreservation any required risk management steps (including washing) should be carried out first, as described in Chapter 4.9.


CHAPTER 4.11.

CATEGORIZATION OF DISEASES AND PATHOGENIC AGENTS

Article 4.11.1.

In 2004, the Research Subcommittee of the International Embryo Transfer Society (IETS) Health and Safety Advisory Committee again reviewed available research and field information on infectious diseases which have been studied regarding the risk of their transmission via in vivo derived embryos. As a result of this review, the IETS has categorised the following diseases and pathogenic agents into four categories.

Please note that this categorisation applies only to in vivo derived embryos.

The following methodology is used by the Research Subcommittee to categorise infectious diseases with regard to the risk of their transmission:

1. Research procedures used to handle and process the embryos will comply with criteria that have been set out by A. Bielanksi and W.C.D. Hare in Appendix A of the IETS Manual.

2. The data used by the Subcommittee to categorise or re-categorise diseases will have been published in peer-reviewed articles in reputable scientific journals. This is to ensure that scientific procedures and results, as well as the interpretation of results, have undergone another level of review.

3. Decisions regarding disease categorisation are based on a consensus judgement which is taken annually by the Subcommittee. The names of members of the Subcommittee who are present when the decisions are made are recorded, as are the names of any others whose opinions were solicited in the decision making process.

4. Questions considered in the decision-making process include the following:
   a) What is the nature of the disease? For example, is the causal agent a uterine pathogen? Does it occur in blood? Does it persist in blood? Do asymptomatic shedders occur? What is the minimum infective dose?
   b) Has the causal agent been found in the ovarian/oviductal/uterine (O.O.U) environment?
   c) Is the causal agent’s presence in the O.O.U environment incidental or is it a consequence of the pathogenesis of the disease?
   d) Is the causal agent’s presence in the O.O.U environment consistent with obtaining viable embryos?
   e) Has the causal agent been found in flushing fluids?
   f) Has the causal agent been found to penetrate or cross the intact zona pellucida (Z.P.)?
   g) Has the causal agent been found to adhere to the ZP?
Annex IX (contd)

h) Is the causal agent removed by washing the embryo?
i) Will special treatments (e.g. with trypsin) remove or inactivate the causal agent?
j) How many embryos have been transferred with or without disease transmission?
k) What is the accumulated evidence for non-transmission of the disease by embryo transfer?
l) What evidence is there that the disease could be transmitted by embryo transfer?
m) Have negative (or positive) results been duplicated by the same or different investigators?
n) Has evidence been accumulated for different animal species as well as for a range of different types and strains of the causal agent?

Article 4.11.2.

Category 1

Category 1 diseases or pathogenic agents are those for which sufficient evidence has accrued to show that the risk of transmission is negligible provided that the embryos are properly handled between collection and transfer according to the IETS Manual.

The following diseases or pathogenic agents are in category 1:
- Aujeszky’s disease (pseudorabies) (swine): trypsin treatment required
- Bluetongue (cattle)
- Bovine spongiform encephalopathy (cattle)
- Brucella abortus (cattle)
- Enzootic bovine leucosis
- Foot and mouth disease (cattle)
- Infectious bovine rhinotracheitis: trypsin treatment required.

Article 4.11.3.

Category 2

Category 2 diseases are those for which substantial evidence has accrued to show that the risk of transmission is negligible provided that the embryos are properly handled between collection and transfer according to the IETS Manual, but for which additional transfers are required to verify existing data.

The following diseases are in category 2:
- Bluetongue (sheep)
- Caprine arthritis/encephalitis
- Classical swine fever (hog cholera)
- Scrapie (sheep).
Article 4.11.4.

Category 3

Category 3 diseases or pathogenic agents are those for which preliminary evidence indicates that the risk of transmission is negligible provided that the embryos are properly handled between collection and transfer according to the IETS Manual, but for which additional in vitro and in vivo experimental data are required to substantiate the preliminary findings.

The following diseases or pathogenic agents are in category 3:

- Bovine immunodeficiency virus
- Bovine spongiform encephalopathy (goats)
- Bovine viral diarrhea virus (cattle)
- Campylobacter fetus (sheep)
- Foot and mouth disease (swine, sheep and goats)
- Haemophilus somnus (cattle)
- Maedi-visna (sheep)
- Mycobacterium paratuberculosis (cattle)
- N. caninum (cattle)
- Ovine pulmonary adenomatosis
- Porcine reproductive and respiratory disease syndrome (PRRS)
- Rinderpest (cattle)
- Swine vesicular disease.

Article 4.11.5.

Category 4

Category 4 diseases or pathogenic agents are those for which studies have been done, or are in progress, that indicate:

1. that no conclusions are yet possible with regard to the level of transmission risk; or
2. the risk of transmission via embryo transfer might not be negligible even if the embryos are properly handled according to the IETS Manual between collection and transfer.
Annex IX (contd)

The following diseases or pathogenic agents are in category 4:

- African swine fever
- Akabane (cattle)
- Bovine anaplasmosis
- Bluetongue (goats)
- Border disease (sheep)
- Bovine herpesvirus 4
- Chlamydia psittaci (cattle, sheep)
- Contagious equine metritis
- Enterovirus (cattle, swine)
- Equine rhinopneumonitis
- Escherichia coli 09:K99 (cattle)
- Leptospira hortorii- serovar hardjobovis (cattle)
- Leptospira sp. (swine)
- Mycobacterium bovis (cattle)
- Mycoplasma spp. (swine)
- Ovine epididymitis (Brucella ovis)
- Parainfluenza 3 virus (cattle)
- Parvovirus (swine)
- Porcine circovirus (type 2) (pigs)
- Scrapie (goats)
- Trichomonas foetus (cattle)
- Ureaplasma/Mycoplasma spp. (cattle, goats)
- Vesicular stomatitis (cattle, swine).

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CHAPTER 4.12.

SOMATIC CELL NUCLEAR TRANSFER IN PRODUCTION LIVESTOCK AND HORSES

Article 4.12.1.

Preface

Following the first meeting of the OIE ad hoc Group on Biotechnology held from 3 to 5 April 2006, the OIE Biological Standards Commission suggested restricting the mandate “to develop recommendations on the animal health risks arising from somatic cell nuclear transfer (SCNT) cloning of production animals, including criteria for assessing the health of embryos and animals derived from such cloning.” The following Articles are a starting point for identifying, characterising and providing a basis for discussion on the animal health risks associated with SCNT cloning technology.

Article 4.12.2.

Overview

At the first meeting of the ad hoc Group on Biotechnology, it was recommended that the Subgroup on Reproductive Animal Biotechnologies should draft recommendations on risk analysis, based on the life-cycle approach, for biotechnology-derived animals. The definition of ‘Reproductive Animal Biotechnology’ was proposed as “the generation of animals through the use of assisted reproductive technologies (ART), which range from artificial insemination through to technologies involving a significant in-vitro component, such as in vitro fertilisation, embryo transfer, embryo splitting and including asexual reproduction such as nuclear transfer”. The following recommendations are restricted to SCNT and are based on a risk analysis approach to biotechnology-derived animals categorised according to the life-cycle approach consisting of: i) embryos, ii) recipients, iii) offspring, and iv) progeny of animal clones.

Article 4.12.3.

Scope

These recommendations address animal health aspects of production animals derived from some reproductive biotechnologies.

Recognising the mandate of the OIE and the suggestion of the OIE Biological Standards Commission, it is the recommendation of the ad hoc Group on Biotechnology to identify risk analysis parameters for animal health and their implication for environmental safety and food and feed safety. These recommendations will focus initially on the scientific basis for the risk assessment aspects, prevention measures and guidance for production livestock and horses derived from SCNT cloning. This is without prejudice to the addition of any relevant issue at a later stage. At present, these recommendations include the following:

- identification of animal health risks and recommendations for management of those risks in embryos, recipients, animal clones and progeny of clones;
- risk and prevention measures related with SCNT cloning technology;
- some welfare issues related to animal health.
Recognising further that the following issues have been discussed or may be addressed by other bodies or instruments, or that they may be addressed at a later stage by the OIE, the document does not address:

- safety and nutritional aspects of food derived from Assisted Reproductive Technologies (ART), for example transgenics (addressed by Codex);
- risks related to the environmental release of animal clones;
- risks related to transgenic animals that have not involved SCNT or other cloning technology;
- non-reproductive animal biotechnologies;
- risks related to animals produced for xenotransplantation or organ donors;
- technologies related to stem cells;
- risks related to aquatic animal health, including fish clones;
- risks related to other terrestrial animals, such as wild mammals and non-mammals, including avian species and insects.

Article 4.12.4.

Background: risk analysis - general principles

1. Risk analysis in general includes hazard identification, risk assessment, risk management and risk communication. The risk assessment is the component of the analysis that estimates the risks associated with a hazard (see Chapter 2.1.). These principles are routinely used by regulators in making decisions about experimental or commercial releases. These analyses can then be used to determine whether the outcomes require management or regulation. Risk management is the process by which risk managers evaluate alternative actions or policies in response to the result(s) of the risk assessment taking into consideration the various social, economic, and legal considerations that form the environment in which such activities occur.

2. For animal diseases, particularly those listed in the Terrestrial Code, there is broad agreement concerning the likely risks and risks can be qualitative or quantitative (see Chapter 2.1.). In disease scenarios it is more likely that a qualitative risk assessment is all that is required. Qualitative assessments do not require mathematical modelling to carry out routine decision-making. Quantitative or semi-quantitative risk assessments assign magnitudes to the risks in numerical (e.g. 1/1,000,000) or descriptive (high/medium/low) terms.

3. In the context of animal cloning, two broad categories of risk assessments are considered: absolute risk assessment and comparative risk assessments. Absolute risk assessments characterise risk independent of a comparator (e.g. the likelihood of an animal transmitting a specific livestock disease). A comparative risk assessment (or relative risk assessment) puts the risk in the context of a comparator.

For example the degree to which an animal produced by one reproductive technology can transmit a particular disease to another animal of the same species compared with the degree to which a similar animal produced by another reproductive technology transmits the same disease to another animal of same species.
4. Regardless of the methodology used, hazard identification is an early step in all science-based risk assessments. In the context of assessing the risks associated with animal cloning (SCNT) and starting with the embryo and moving on through animal clone development and subsequent progeny, it is important to be clear at this juncture that only a comparative semi-quantitative risk assessment can be completed. A systematic, absolute, quantitative risk assessment of potential risks is difficult, due to the relative newness of the technology, and the variability in outcomes among laboratories and species cloned. Furthermore, with the technology of SCNT there is no introduced hazard from the insertion of novel genes (which may potentially happen in transgenesis). Thus, to analyse what factors contribute to animal health risks, the existing baseline must be analysed.

5. In short, the specific points where the risk assessment needs to be focused need to be identified. As illustrated in the accompanying diagram - the focus is to look at the basics of creating an embryo - using current terminology, starting from the selection of donor of oocyte and the cells to the creation of an embryo by the cloning methodology. The second phase will focus on the recipient of the embryo clone and the animal health and care considerations for the animals. The actual embryo clone that is born as an offspring is the third part of the paradigm that needs clear recommendations for assessment, and the next generation, either the progeny of the animal clone (which is a result of normal sexual reproduction) or animals produced by recloning (clones of clones) is the fourth and final stage.

**Article 4.12.5.**

**Managing animal health risks associated with embryos**

Embryo production by in vitro techniques has been applied for many years. Although the additional steps involved in cloning add a new dimension to this procedure, many of the risks associated with SCNT have previously been identified for established ART animal reproductive biotechnologies (see Chapter 4.8.). An analysis of SCNT methodology allows the procedural details to be categorised into:

a) Oocytes (obtained from the abattoir, recovered from trans-vaginal ultrasound-guided procedures or by laparotomy procedures)

   Ovaries which are collected at an abattoir should be collected, transported and processed according to the recommendations laid down in Chapter 4.8.

   The primary risks are associated with the health status of the animal from which the oocytes are harvested and the quality of the oocytes.

b) Donor cells (cells obtained from animals chosen to be cloned – by biopsy, harvesting at slaughter or after death)

   Currently there are no specific new risks identified with SCNT cloning. There is a proposed risk related to activation of endogenous retroviruses during cell transfer procedures, however, this may be more theoretical than practical. In some current experimental procedures, the donor cell may be treated with chemicals to modify its composition, for example cell cycle inhibitors or chromatin modifiers.

c) In vitro culture of reconstructed embryos (procedure used to fuse the donor and recipient material and to culture the reconstructed embryo)

d) Risks associated with the method of fusing donor cells with enucleated recipient oocytes and with culture conditions.

In addition, the practitioner should ensure that the clone pregnancy is compatible to the surrogate dam’s breed, anatomy and physiology.
SOMATIC CELL NUCLEAR TRANSFER

Oocyte donor

Mature Oocyte

Removing the nucleus from Oocyte - Enucleation

Enucleated Oocyte

Transfer of donor cell into the enucleated oocyte

Fusion of the cell with the enucleated oocyte

SCNT Embryo

Transfer of the cloned embryo in the recipient

Recipient

Birth of a SCNT

Pregnancy

SCNT Clone

Progeny

Replication by cloning
Or by sexual reproduction

© HPSK
1. **Oocytes**

The laboratory or the producer should establish a detailed record of ovaries – their origin, health of the animal from which the ovaries are obtained, details of any systemic lesion on the animal and proper herd data. This is particularly useful where the pooling of ovaries may provide cross-contamination of ovarian tissue.

Follicular fluids may carry various infectious agents like bovine viral diarrhoea virus (BVDV) and can contaminate pooled follicular fluid from healthy animals. Furthermore, the technique for collecting oocytes, such as aspiration or slicing of the ovarian follicles, determines the extent of blood contamination or extraneous material. A representative sample to demonstrate the absence of infectious biological material should be done with each pooled batch.

Oocytes are matured as cumulus oocyte complexes (COCs) and then matured in most instances in the culture/maturation media. Care and efforts should be taken to carefully select and mature the oocytes from the pools that are morphologically good; also the media used should have been quality tested. Use of serum or protein components from an undefined or untested source should be avoided. Addition of proper and safe antibiotics in the culture media to control opportunistic bacteria should be encouraged.

Use of proper sanitary and disinfection procedures is of utmost importance and should be emphasised in any in vitro fertilisation (IVF) laboratory. Proper handling and following sanitary protocols during the maturation and further culture of embryos should be encouraged.

2. **Donor cells**

In order to minimise risks:

- Donor cells should be properly harvested from the animal and cultured under proper sanitary conditions using good laboratory practices.

- When applicable, the passaging of the cells used for the cloning procedure should be documented and at different stage sampling may be warranted to look at the chromosomal component of the cell lines. If possible, procedures should be in place for regular sampling of the cells for morphological and other characteristics.

- Master cell lines (to be used for cloning at a later stage) should be stored under conditions found to be optimal for maintaining viability. Freedom from extraneous agents should be established by testing for bacteria, fungi, mycoplasmas or viruses, using appropriate tests (see Manual of the International Embryo Transfer Society [IETS]).

3. **Cloning procedures/reconstruction**

The cloning procedure that employs the use of chemicals or other reagents should be carefully evaluated, in terms of the quality of embryos and overall efficiency.

During the fusion of recipient and donor material by chemical or physical means care and control should be employed. The optimisation of the procedure based on the laboratory protocols or published reports should be determined to avoid early embryonic mortalities.
Annex X (contd)

If co-culture of the cell is used for the culture procedure after reconstruction of embryos, proper screening of the co-culture cells should be done. A sample of each batch may be tested for the bacterial, fungal, mycoplasmal or viral component.

Embryos should be cultured and harvested for an appropriate time and stage to transfer them or to cryo-preserve them for later use. Proper procedures based on the international standards (IETS Codes of Practice) for washing and preservation of the embryos should be followed.

Care should be taken with regard to grading the embryos before transfer (see Chapters 4.7. and 4.8.).

Article 4.12.6.

Managing animal health risks related to the recipients (surrogate dams)

1. Animal health risks to the surrogate dams

Currently, when compared with in vitro produced embryos, SCNT has a higher rate of pregnancy failure and, in some species, placental abnormalities. Loss due to defects in the embryo or failure to implant in the uterus of the surrogate dam does not pose a hazard to the dam. Rather, the surrogate dam simply resorbs any embryonic tissue and returns to cycling. Mid- and late-term spontaneous abortions may be hazardous to surrogates if they are unable to expel the fetus and its associated membranes. Most abortions in natural service and artificial insemination (AI) pregnancies in cattle remain undiagnosed due to the expense of laboratory work and the low profit margin in both the beef and dairy industry. Producers and veterinarians become concerned when the rate of abortion exceeds 3–5% in a herd. The same potential impact of external influences should be considered with pregnancy evaluation with SCNT and other reproductive technologies. Disease, under-nutrition, and severe environmental conditions are stressors known to interfere with animal fertility and embryo survival. Under these circumstances, the risk to the pregnancy is directly related to stress factors and not to the technology used.

To date, a species-specific effect has been seen. Abnormalities in clones may result from incomplete reprogramming of the donor nucleus. Epigenetic reprogramming occurs at different times in embryos in different species. Many of the abnormalities reported in cattle and sheep pregnancies have not been noted in goats or swine carrying SCNT clones. The amount of in vitro manipulation of an embryo inversely correlates to the chances for successful pregnancy outcomes. This has been observed in both SCNT embryos and in vitro produced fertilised embryos. Unlike other forms of other reproductive technologies SCNT pregnancy losses occur at all stages of gestation in cattle. Clone pregnancies have been lost during the second and third trimesters and have been accompanied by reports of hydrops, enlarged umbilicus, and abnormal placentation.

2. Animal health risks posed by the surrogate dam to the clone embryos

No new animal health risks have been identified for the developing clone fetus from the surrogate dam compared with conventional pregnancies. The latter include vertically transmitted diseases and abnormalities due to metabolic or physiological stress.

With respect to the animal health risks associated with the surrogate dam, it is difficult to document the relative frequency of early stage losses of SCNT embryos compared with early stage losses of other pregnancies as these abortions are not typically diagnosed with other reproductive technologies. Additionally, external stressors will similarly impact SCNT pregnancies.
Veterinarians should monitor the progress of pregnancy as the common gestational anomalies seen in other assisted reproductive technologies may be exhibited and diagnosed during the physical examination. A database of commonly encountered problems in clone pregnancies would be useful if available to animal health experts.

- Care should be taken to assess the general health of the recipient dam before selection to carry the embryo clones. The general health status of the recipient should be determined in terms of freedom from infection and disease, proper vaccination and follow-up, and, if applicable, proof of earlier uneventful pregnancies, absence of birthing problems, and proper post-pregnancy recovery.

- Pregnancy loss is greatest with SCNT embryos prior to 60 days’ gestation in cattle. This is similar to the pattern seen with other reproductive technologies. However, in clones, high pregnancy losses during this time of placental formation (between 45–60 days) suggest that embryonic death may be a consequence of faulty placentation. Abnormal placentation may lead to a build up of wastes in the fetus and associated membranes, or inadequate transfer of nutrients and oxygen from the dam to the fetus. Care should be taken to monitor the recipient dam during pregnancy. Once the pregnancy is established and confirmed, regular veterinary assessments and monitoring of animal health status is desirable up to the birth of the offspring.

- To ensure that the recipient is pregnant and to monitor its health during the first trimester, it is useful to perform ultrasonographic assessments, determine hormonal profiles and assess the general physiological parameters. Based on these profiles, proper attention should be paid to aid in the proper establishment of pregnancy by providing proper husbandry conditions and nutrition.

- The animals should be observed carefully for the signs of labour nearing the time of birth. In some species, one of the more common problems is uterine inertia and the absence of contractions. The absence of contractions may result in prolonged pregnancies with associated sequelae that may require assistance with deliveries.

- A surgical intervention should be decided and should be available for the near term animal if the situation so warrants. Proper procedures should be employed to ascertain the proper handling of the offspring and the surrogate dam.

- Health concerns may arise as a result of surgical procedures, excessive traction, or other complications such as retained fetal membranes. In these cases post-partum care may be necessary.

3. Managing animal health risks of animal clones

The health problems of individual clones can be observed in utero and post-partum. These appear to be the same as observed in other ART assisted reproductive technologies, but they may be more common in clones. It is important to determine whether the abnormalities are of genetic or epigenetic origin. Large offspring syndrome (LOS), probably in relation to placental abnormalities rather than fetal abnormalities, have been particularly observed in cloned cattle and sheep following suboptimal in vitro handling. These abnormalities are becoming less frequent in small ruminants.

- Appropriate husbandry practices are important to the health of animal clones. Care should be taken to provide colostrums and a clean and hygienic environment, supervision for the first few weeks after birth should be practiced.
- The animal clones must be checked routinely for the most common phenotypic anomalies, such as atresia ani, umbilical hernia, flexor muscle contractions, respiratory or cardiac insufficiency, and failure to suckle. This will allow proper treatment and care of the newborn and increase the survival of the young one.

- To consolidate current understanding of the health status of animal clones, a comprehensive veterinary examination should be performed to monitor the progress of the clone, as unexplained fatalities or fatalities arising from systemic complications have been reported. It is encouraged to follow the health profile of the animals to at least the reproductive maturity stage, and to record the ability to reproduce (fertility index).

- Animal welfare concerns ranging from LOS to serious abnormalities are notable in the debates pertaining to cloning technology. Proper research and peer-reviewed data should be generated. The animal clones should undergo species-specific basic welfare assessments. If welfare concerns are detected at initial screening, a more extensive characterisation of that phenotype should be performed to document the animal welfare concerns.

- Proper monitoring of the animal population during different stages of life from birth to puberty should be documented to address and validate the genomic potential of the animal clones.

4. Managing animal health risks related to sexually reproduced progeny of clones

Presently there is no evidence of an increased health risk if sexual reproduction is used for obtaining progeny. Some data indicate that the reprogramming errors during the cloning process may actually be corrected during the natural mating and reproduction process:

a) Characterisation of the health profile, including health status and data on animal welfare, would consolidate the knowledge of sexually reproduced progeny.

b) Monitoring the reproductive performance of sexually reproduced progeny of clones would be useful to assess their reproductive capacity in comparison with their conventional counterparts.

5. Managing animal health risks associated with re-cloning/ clones of clones

Information on recloning is only beginning to appear. It is therefore necessary to follow the approach below:

a) The health profile (health status and data on animal welfare) should be characterised to consolidate the knowledge.

b) The reproductive performance of clones of clones should be monitored to assess the capacity of the animals to perform in comparison with their conventional counterparts.
Article 4.12.7.

Review

The goal of this Chapter is to provide a scientific basis and recommendations on animal health and welfare risks to animals involved in SCNT cloning compared with other ART assisted reproductive technologies. These recommendations will focus initially on the scientific basis for the risk assessment aspects, prevention measures and guidance for production livestock and horses, derived from SCNT cloning and should be reviewed in light of new scientific information.

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CHAPTER 5.1.
GENERAL OBLIGATIONS RELATED TO CERTIFICATION

Article 5.1.1.

Safety of international trade in animals and animal products depends on a combination of factors which should be taken into account to ensure unimpeded trade, without incurring unacceptable risks to human and animal health.

Because of differences between countries in their animal health situations, various options are offered by the Terrestrial Code. The 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 sanitary aspects of international trade, Veterinary Authorities of OIE Members should base their import requirements on the OIE standards.

These requirements should be included in the model certificates approved by the OIE which are included from Chapters 5.10 to 5.12 of the Terrestrial Code.

Certification requirements should be exact and concise, and should clearly convey the wishes of the importing country. For this purpose, prior consultation between Veterinary Authorities of importing and exporting countries may be necessary. It enables the setting out of the exact requirements so that the signing veterinarian can, if necessary, be given a note of guidance explaining the understanding between the Veterinary Authorities involved.

When officials of a Veterinary Authority wish to visit another country for matters of professional interest to the Veterinary Authority of the other country, the latter should be informed.

Article 5.1.2.

Responsibilities of the importing country

1. The import requirements included in the international veterinary certificate should assure that commodities introduced into the importing country comply with the OIE standards. Importing countries should restrict their requirements to those necessary to achieve the national appropriate level of protection. If these are stricter than the OIE standards, they should be based on an import risk analysis.

2. The international veterinary certificate should not include requirements for the exclusion of pathogens or animal diseases which 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 specific pathogen or disease should not require a higher level of protection than that provided by measures applied as part of the official control programme operating within the importing country.

3. The international veterinary certificate should not include measures against pathogens or diseases which are not OIE listed, unless the importing country has demonstrated through import risk analysis, carried out in accordance with Section 2, that the pathogen or disease poses a significant risk to the importing country.

4. The transmission by the Veterinary Authority of certificates or the communication of import requirements to persons other than the Veterinary Authority of another country, necessitates that copies of these documents are also sent to the Veterinary Authority. This important procedure avoids delays and difficulties which may arise between traders and Veterinary Authorities when the authenticity of the certificates or permits is not established.
Annex XI (contd)

This information is the responsibility of Veterinary Authorities. 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 the Veterinary Authority.

5. Situations may arise which result in changes to the consignee, identification of the means of transportation, or border post after a certificate is issued. Because these do not change the animal or public health status of the consignment, 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 animal health situation and national animal health information systems to determine whether that country is free or has free zones or compartments free of listed diseases, including the regulations and procedures in force to maintain its free status;

   b) regular and prompt information on the occurrence of notifiable diseases;

   c) details of the country's ability to apply measures to control and prevent the relevant listed diseases;

   d) information on the structure of the Veterinary Services and the authority which they exercise according to Chapters 3.1. and 3.2.;

   e) technical information, particularly on biological tests and vaccines applied in all or part of the national territory.

2. Veterinary Authorities of exporting countries should:

   a) have official procedures for authorisation of certifying veterinarians, defining their functions and duties as well as conditions covering possible suspension and termination of the appointment;

   b) ensure that the relevant instructions and training are provided to certifying veterinarians;

   c) monitor the activities of the certifying veterinarians to verify their integrity and impartiality.

3. The Head of the Veterinary Service Authority of the exporting country is ultimately accountable for veterinary 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 the recognised incubation periods of the various diseases subsequent to an export taking place, the Veterinary Authority becomes aware of the appearance or reappearance of a disease which has been specifically included in the international veterinary certificate, there is an obligation for this 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. Equally, if a disease condition appears in imported commodities within a time period after importation consistent with the recognised incubation period of the disease, the Veterinary Authority of the exporting country should be informed so as to enable an investigation to be made, since this may be the first available information on the occurrence of the disease in a previously free herd. The Veterinary Authority of the importing country should be informed of the result of the investigation since the source of infection may not be in the exporting country.

3. In case of suspicion, on reasonable grounds, that an official certificate may be fraudulent, the Veterinary Authority of the importing country and exporting country should conduct an investigation. Consideration should also be given to notifying any third country(ies) that may have been implicated. All associated consignments should be kept under official control, pending the outcome of the investigation. The Veterinary Authorities of all countries involved should fully cooperate with the investigation. If the certificate is found to be fraudulent, every effort should be made to identify those responsible so that appropriate action can be taken according to the relevant legislation.
CHAPTER 5.2.

CERTIFICATION PROCEDURES

Article 5.2.1.

Protection of the professional integrity of the certifying veterinarian

Certification should be based on the highest possible ethical standards, the most important of which is that the professional integrity of the certifying veterinarian must be respected and safeguarded according to Chapters 3.1. and 3.2.

It is essential not to include in the requirements additional specific matters which cannot be accurately and honestly signed by a veterinarian. For example, these requirements should not include certification of an area as being free from non-notifiable diseases the occurrence of which the signing veterinarian is not necessarily informed about. Equally, to ask certification for events which will take place after the document is signed is unacceptable when these events are not under the direct control and supervision of the signing veterinarian.

Certification of freedom from diseases based on purely clinical freedom and herd history is of limited value. This is also true of diseases for which there is no specific diagnostic test, or the value of the test as a diagnostic aid is limited.

The note of guidance referred to in Article 5.1.1. is not only to inform the signing veterinarian but also to safeguard professional integrity.

Article 5.2.2.

Certifying veterinarians

Certifying veterinarians should:

1. be authorised by the Veterinary Authority of the exporting country to sign international veterinary 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 competent party;

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 veterinarian should be in possession of that documentation before signing;

4. have no conflict of interest in the commercial aspects of the animals or animal products being certified and be independent from the commercial parties.

Article 5.2.3.

Preparation of international veterinary certificates

Certificates should be drawn up in accordance with the following principles:
Annex XI (contd)

1. Certificates should be designed so as to minimize the potential for fraud including use of a unique identification number, or other appropriate means to ensure security. Paper certificates should bear the official identifier of the issuing Veterinary 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. They should be written in terms that are as simple, unambiguous and easy to understand as possible, without losing their legal meaning.

3. If so required, they 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 veterinarian.

4. They should require appropriate identification of animals and animal products except where this is impractical (e.g. day-old birds).

5. They should not require a veterinarian to certify matters that are outside his/ her knowledge or which he/ she cannot ascertain and verify.

6. Where appropriate, they should be accompanied, when presented to the certifying veterinarian, by notes of guidance indicating the extent of enquiries, tests or examinations expected to be carried out before the certificate is signed.

7. Their text should not be amended except by deletions which must be signed and stamped by the certifying veterinarian.

8. The signature and stamp must 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. Replacement certificates may be issued by a Veterinary Authority to replace certificates that have been, for example, lost, damaged, contain errors, or where the original information is no longer correct. These duplicates replacements should be provided by the issuing authority and these must 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.

10. Only original certificates are acceptable.

Article 5.2.4.

Electronic certification

1. Certification may be provided by electronic documentation sent directly from the Veterinary Authority of the exporting country to the Veterinary Authority of the importing country. Such systems also normally provide an interface with the commercial organisation marketing the commodity for provision of information to the certifying authority. The certifying veterinarian must have access to all information such as laboratory results and animal identification data.

2. Electronic certificates may be in a different format but should carry the same information as conventional paper certificates.
3. The Veterinary Authority must have in place systems for the security of electronic certificates against access by unauthorised persons or organisations.

4. The certifying veterinarian must be officially responsible for the secure use of his/her electronic signature.

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CHAPTER 6.1.

THE ROLE OF THE VETERINARY SERVICES
IN FOOD SAFETY

Article 6.1.1.

Purpose

The purpose of this Chapter is to provide guidance to OIE Members in regard to the role and responsibilities of the Veterinary Services in food safety, to assist them in meeting the food safety objectives laid down in national legislations and the requirements of importing countries.

Article 6.1.2.

Background

Historically, the Veterinary Services were set up to control livestock diseases at the farm level. There was an emphasis on prevention and control of the major epizootic diseases of livestock and of diseases that could affect man (zoonotic diseases). As countries begin to bring the serious diseases under control, the scope of official animal health services normally increases to address production diseases of livestock, where control leads to more efficient production and/or better quality animal products.

The role of the Veterinary Services has traditionally extended from the farm to the slaughterhouse, where veterinarians have a dual responsibility – epidemiological surveillance of animal diseases and ensuring the safety and suitability of meat. The education and training of veterinarians, which includes both animal health (including zoonoses) and food hygiene components, makes them uniquely equipped to play a central role in ensuring food safety, especially the safety of foods of animal origin. As described below, in addition to veterinarians, several other professional groups are involved in supporting integrated food safety approaches throughout the food chain. In many countries the role of the Veterinary Services has been extended to include subsequent stages of the food chain in the “farm to fork” continuum.

Article 6.1.3.

Approaches to food safety

1. The concept of the food production continuum

Food safety and quality are best assured by an integrated, multidisciplinary approach, considering the whole of the food chain. Eliminating or controlling food hazards at source, i.e. a preventive approach, is more effective in reducing or eliminating the risk of unwanted health effects than relying on control of the final product, traditionally applied via a final ‘quality check’ approach. Approaches to food safety have evolved in recent decades, from traditional controls based on good practices (Good Agricultural Practice, Good Hygienic Practice, etc.), via more targeted food safety systems based on hazard analysis and critical control points (HACCP) to risk-based approaches using food safety risk analysis.

2. Risk-based management systems

The development of risk-based systems has been heavily influenced by the World Trade Organization Agreement on the Application of Sanitary and Phytosanitary Measures (“SPS Agreement”). This Agreement stipulates that signatories shall ensure that their sanitary and phytosanitary measures are based on an assessment of the risks to human, animal or plant life or health, taking into account risk assessment techniques developed by relevant international organizations. Risk assessment, the scientific component of risk analysis, should be functionally separated from risk management to avoid interference from economic, political or other interests.
The SPS Agreement specifically recognises as the international benchmarks the standards developed by the OIE for animal health and zoonoses and by the Codex Alimentarius Commission for food safety. In recent decades there has also been a trend towards a redefinition of responsibilities. The traditional approach, whereby food operators were primarily held responsible for food quality while regulatory agencies were charged with assuring food safety, has been replaced by more sophisticated systems that give food operators primary responsibility for both the quality and the safety of the foods they place on the market. The role of the supervisory authorities is to analyse scientific information as a basis to develop appropriate food safety standards (both processing and end product standards) and monitoring verification inspections to ensure that the control systems used by food operators are appropriate, validated, effective and operated in such a way that the standards are met. In the event of non-compliance, regulatory agencies are responsible to ensure that appropriate corrective actions are taken and sanctions are applied.

The Veterinary Services play an essential role in the application of the risk analysis process and the implementation of risk-based recommendations for regulatory systems, including the extent and nature of veterinary involvement in food safety activities throughout the food chain, as outlined above. Each country should establish its health protection objectives, for animal health and public health, through consultation with stakeholders (especially livestock producers, processors and consumers) in accordance with the social, economic, cultural, religious and political contexts of the country. These objectives should be put into effect through national legislation and policies and steps taken to raise awareness of them both within the country and to trading partners.

3. Functions of Veterinary Services

The Veterinary Services contribute to the achievement of these objectives through the direct performance of some veterinary tasks and through the auditing of animal and public health activities conducted by other government agencies, private sector veterinarians and other stakeholders. In addition to veterinarians, several other professional groups are involved in ensuring food safety throughout the food chain, including analysts, epidemiologists, food technologists, human and environmental health professionals, microbiologists and toxicologists. Irrespective of the roles assigned to the different professional groups and stakeholders by the administrative system in the country, close cooperation and effective communication between all involved is imperative to achieve the best results from the combined resources. Where veterinary or other professional tasks are delegated to individuals or enterprises outside the Veterinary Authority, clear information on regulatory requirements and a system of checks should be established to monitor and verify performance of the delegated activities. The Veterinary Authority retains the final responsibility for satisfactory performance of delegated activities.

4. At the farm level

Through their presence on farms and appropriate collaboration with farmers, the Veterinary Services play a key role in ensuring that animals are kept under hygienic conditions and in the early detection, surveillance and treatment of animal diseases, including conditions of public health significance. The Veterinary Services may also provide livestock producers with information, advice and training on how to avoid, eliminate or control food safety hazards (e.g. drug and pesticide residues, mycotoxins and environmental contaminants) in primary production, including through animal feed. Producers’ organisations, particularly those with veterinary advisors, are in a good position to provide awareness and training as they are regularly in contact with farmers and are well placed to understand their priorities. Technical support from the Veterinary Services is important and both private veterinarians and employees of the Veterinary Authority can assist. The Veterinary Services play a central role in ensuring the responsible and prudent use of biological products and veterinary drugs, including antimicrobials, in animal husbandry. This helps to minimise the risk of developing antimicrobial resistance and unsafe levels of veterinary drug residues in foods of animal origin. Chapters 6.5. to 6.8. of the Terrestrial Code contain recommendations on the use of antimicrobials.
5. Meat inspection

Slaughterhouse inspection of live animals (ante-mortem) and their carcasses (post-mortem) plays a key role in both the surveillance network for animal diseases and zoonoses and ensuring the safety and suitability of meat and by-products for their intended uses. Control and/or reduction of biological hazards of animal and public health importance by ante- and post-mortem meat inspection is a core responsibility of the Veterinary Services and they should have primary responsibility for the development of relevant inspection programmes.

Wherever practicable, inspection procedures should be risk-based. Management systems should reflect international standards and address the significant hazards to both human and animal health in the livestock being slaughtered. The Codex Alimentarius Code of Hygienic Practice for Meat (CHPM) constitutes the primary international standard for meat hygiene and incorporates a risk-based approach to application of sanitary measures throughout the meat production chain. Chapter 6.2. of the Terrestrial Code contains recommendations for the control of biological hazards of animal health and public health importance through ante- and post-mortem meat inspection, which complement the CHPM.

Traditionally, the primary focus of the Terrestrial Code was on global animal health protection and transparency. Under its current mandate, the OIE also addresses animal production food safety risks. The Terrestrial Code includes several standards and recommendations aimed at protecting public health (such as Chapter 6.2. on the Control of Biological Hazards of Animal Health and Public Health Importance through Ante- and Post- Mortem Meat Inspection) and work is underway developing new standards to prevent the contamination of animal products by Salmonella spp. and Campylobacter spp. The OIE and Codex collaborate closely in the development of standards to ensure seamless coverage of the entire food production continuum. The recommendations of the OIE and the Codex Alimentarius Commission on the production and safety of animal commodities should be read in conjunction.

The Veterinary Authority should provide for flexibility in the delivery of the meat inspection service. Countries may adopt different administrative models, involving degrees of delegation to officially recognised competent bodies operating under the supervision and control of the Veterinary Authority. If personnel from the private sector are used to carry out ante- and post-mortem inspection activities under the overall supervision and responsibility of the Veterinary Authority, the Veterinary Authority should specify the competency requirements for all such persons and verify their performance. To ensure the effective implementation of ante- and post-mortem inspection procedures, the Veterinary Authority should have in place systems for the monitoring of these procedures and the exchange of information gained. Animal identification and animal traceability systems should be integrated in order to be able to trace slaughtered animals back to their place of origin, and products derived from them forward in the meat production chain.

6. Certification of animal products for international trade

Another important role of the Veterinary Services is to ensure that health certification for international trading partners attesting that exported products meet both trade complies with animal health and food safety standards. Certification in relation to animal diseases, including zoonoses, and meat hygiene should be the responsibility of the Veterinary Authority. Certification may be provided by other professions (a sanitary certificate) in connection with food processing and hygiene (e.g. pasteurisation of dairy products) and conformance with product quality standards.
Annex XII (contd)

7. The roles of the Veterinary Services

Most reported outbreaks of foodborne disease are due to contamination of foods with zoonotic agents, often during primary production. The Veterinary Services play a key role in the investigation of such outbreaks all the way back to the farm and in formulating and implementing remedial measures once the source of the outbreak has been identified. This work should be carried out in close collaboration with human and environmental health professionals, analysts, epidemiologists, food producers, processors and traders and others involved.

In addition to the roles mentioned above, veterinarians are well equipped to assume important roles in ensuring food safety in other parts of the food chain, for example through the application of HACCP-based controls and other quality assurance systems during food processing and distribution. The Veterinary Services also play an important role in raising the awareness of food producers, processors and other stakeholders of the measures required to assure food safety.

8. Optimising the contribution of the Veterinary Services to food safety

In order for Veterinary Services to make the best possible contribution to food safety, it is important that the education and training of veterinarians in the roles outlined in this Chapter meets high standards and that there are national programmes for ongoing and comprehensive professional development. The Veterinary Services should comply with the OIE fundamental principles of quality given in Chapter 3.1. of the Terrestrial Code. Recommendations for the evaluation of Veterinary Services are provided in Chapter 3.2. of the Terrestrial Code and in the OIE Tool for the Evaluation of Performance of Veterinary Services.

There should be a clear and well documented assignment of responsibilities and chain of command within the Veterinary Services. The national Competent Authority should provide an appropriate institutional environment to allow the Veterinary Services to develop and implement the necessary policies and standards and adequate resources for them to carry out their tasks in a sustainable manner. In developing and implementing policies and programmes for food safety, the Veterinary Authority should collaborate with other responsible agencies to ensure that food safety risks are addressed in a coordinated manner.
CHAPTER X.X.

PREVENTION, DETECTION AND CONTROL OF SALMONELLA IN POULTRY

Article X.X.1.

Introduction

This Chapter provides recommendations on the prevention, detection and control of Salmonella in poultry.

Salmonellosis is one of the most common foodborne bacterial diseases in the world. The great majority of Salmonella infections in humans are foodborne with Salmonella Enteritidis and Salmonella Typhimurium accounting for a major part of the problem. Salmonella serotypes and prevalence may vary considerably between localities, districts, regions and countries and therefore, surveillance and identification of the prevalent Salmonella serotypes in humans and poultry should be carried out in order to develop a control programme for the area.

In most food animal species, Salmonella can establish a clinically inapparent infection of variable duration, which is significant as a potential zoonosis. Such animals may be important in relation to the spread of infection between flocks and as causes of human foodborne infection. In the latter case, this can occur when meat and eggs, or their products, enter the food chain thus producing contaminated food.

Article X.X.2.

Purpose and scope

This Chapter deals with methods for on farm prevention, detection and control of Salmonella in poultry, and complements the Codex Alimentarius Code of Hygiene Practice for Meat (CAC/ RCP 58-2005) and Code of Hygienic Practice for Eggs and Egg Products (CAC/ RCP 15-1976 Revision 2007). A pathogen reduction strategy at the farm level is seen as the first step in a continuum that will assist in reducing the presence of foodborne pathogens in eggs and meat.

Hygiene and biosecurity procedures to be implemented in poultry flocks and hatcheries are described in Chapter 6.3. Hygiene and Biosecurity Procedures in Poultry Production.

The recommendations presented in this Chapter are relevant to the control of all Salmonella with special attention to S. Enteritidis and S. Typhimurium, as these are common Salmonella serotypes in many countries. It should be noted that the epidemiology of animal and human salmonellosis in a particular locality, district, region or country is important for effective control of Salmonella.

Article X.X.3.

Definitions (for this Chapter only)

Breeders

means poultry destined for the production of fertile eggs for incubation for the purpose of producing day-old chicks.
Annex XIII (contd)

**Competitive exclusion**
means the administration of defined or undefined bacterial flora to poultry to prevent gut colonisation by enteropathogens, including *Salmonella*.

**Culling**
means the depopulation of a flock before the end of its normal production period.

**Layers**
means poultry during the period of laying eggs for human consumption.

**Poultry**
means all domesticated birds, including backyard poultry, used for the production of meat or eggs for consumption, for the production of other commercial products, for restocking supplies of game, or for breeding these categories of birds, as well as fighting cocks used for any purpose. Birds that are kept in captivity for any reason other than those reasons referred to in the preceding paragraph, including those that are kept for shows, races, exhibitions, competitions or for breeding or selling these categories of birds as well as pet birds, are not considered to be poultry.

**Article X.X.4.**

**Surveillance of poultry flocks for Salmonella**

Where justified by risk assessment, surveillance should be carried out to identify infected flocks in order to take measures that will reduce the prevalence in poultry and the risk of transmission of *Salmonella* to humans. Sampling methods, frequency and type of samples required should be determined by the Veterinary Services based on risk assessment. Microbiological testing is preferred to serological testing because of its higher sensitivity in broilers flocks and higher specificity in breeders and layer flocks. In the framework of regulatory programmes for the control of *Salmonella* in poultry and salmonellosis in humans, confirmatory testing may be required to ensure that decisions are soundly based.

**Sampling**

1. **Available methods for sampling**

   Drag swabs: sampling is done by dragging swabs throughout the poultry building.

   Boot swabs: sampling is done by walking throughout the poultry building with absorbent material placed over the footwear of the sampler.

   Faecal samples: multiple fresh faecal/caecal samples collected from different areas in the poultry building.

   Meconium, chick box papers, dead in shell and culled chicks at the hatchery.

   Hatchery samples: throughout the hatchery, including the inner liner of the incubators.

   Additional sampling of equipment and surfaces may be performed to increase sensitivity.

2. **Sample size**

   Refer to the Terrestrial Manual.
3. Laboratory methods

Refer to the Terrestrial Manual.

4. Time and frequency of testing

Time and frequency of sampling for each poultry type are listed below:

a) Breeders and hatcheries

i) Breeder flocks before lay

- Before the end of the first week of life when the status of the breeding farm and the hatchery is not known or does not comply with this Chapter.
- Within the four weeks before being moved to another house, or before going into production if the animals will remain in the same house for the production period.
- One or more times during the growing period if there is a culling policy in place. The frequency would be determined on commercial considerations.

ii) Breeder flocks in lay

- At least at monthly intervals during the laying period.
- Additional testing should be determined by the Veterinary Services.

iii) Hatcheries

- Testing hatcheries may complement farm testing.
- The minimal frequency should be determined by the Veterinary Services.

b) Poultry for the production of eggs for human consumption

i) Flocks grown to be layers

- Before the end of the first week of life when the status of the breeding farm and the hatchery is not known or does not comply with this Chapter.
- Within the four weeks before being moved to another house, or before going into production if the animals will remain in the same house for the production period.
- One or more times during the growing period if there is a culling policy in place. The frequency would be determined on commercial considerations.

ii) Layer flocks

- At expected peak of lay for each production cycle (the period of time in the laying cycle when the production of the flock is highest).
- One or more times if there is a culling policy in place or if eggs are diverted to processing for the inactivation of the pathogen. The minimal frequency should be determined by the Veterinary Services.
Annex XIII (contd)

c) Poultry for the production of meat

i) Flocks should be sampled at least once.

ii) Where sampling occurs on farms and where there is a long period (2 weeks or more) between thinning and final depopulation further testing should be considered.

iii) Where sampling occurs on farms, flocks should be sampled as late as possible before the first birds are transported to the slaughter house. Where this is done to allow for the implementation of control measures during processing, this must be done at a time that ensures the results are available before slaughter.

Whether sampling occurs on the farm or at the processing plant, there should be an integrated system in place which allows for investigation of the source of positive flocks.

d) Empty building testing

i) Bacteriological monitoring of the efficacy of disinfection procedures is recommended when Salmonella have been detected in the previous flock.

As appropriate, sampling of equipment and surfaces as well as boot swabs or drag swabs of the empty building after depopulation, cleaning and disinfection.

Results from surveillance may lead to the implementation of additional prevention and control measures to reduce the risk of transmission of Salmonella to humans:

a) In breeders, control measures may be implemented to reduce the transmission of Salmonella to the next generation, especially for trans-ovarian transmitted serotypes such as S. Enteriditis.

b) In layer flocks control measures will reduce and may eliminate contamination of eggs with Salmonella.

c) In poultry for meat production, control measures may be implemented at slaughter or further down the food chain.

Article X.X.5.

Prevention and Control measures

Salmonella prevention and control be achieved by adopting Good Agricultural Practices and Hazard Analysis Critical Control Point (HACCP), and general measures detailed in Chapter 6.3. Hygiene and Biosecurity Procedures in Poultry Production, in combination with the following additional measures, where appropriate. No single measure used alone will achieve effective Salmonella control.

Additional prevention and control measures include: vaccination, competitive exclusion, flock culling, organic acids and product diversion to processing.
Antimicrobials should not be used to control infection with Salmonella in poultry because the effectiveness of the treatment is limited, may mask the infection at sampling, has the potential to produce residues in meat and eggs and can contribute to the development of antimicrobial resistance. Antimicrobials may also reduce normal flora in the gut and increase the likelihood of colonisation with Salmonella. In special circumstances antimicrobials may be used to salvage animals with high genetic value.

1. Day old chicks used to stock a poultry house should be obtained from breeding flocks and hatcheries that are free from at least S. Enteritidis and S. Typhimurium and have been monitored according to this Chapter.

2. Layer and breeder flocks should be stocked from flocks that are free from at least S. Enteritidis and S. Typhimurium and have been monitored according to this Chapter.

3. Feed contamination with Salmonella is known to be a source of infection for poultry. Therefore, it is recommended to monitor the Salmonella status of poultry feed, and if found positive to take corrective measures. The use of heat treated feeds or feeds subjected to other bacteriostatic or bactericidal treatment (e.g. organic acids) is recommended. Feed should be stored in clean closed containers to prevent access by wild birds and rodents. Spilled feed should be cleaned up immediately to remove attractants for wild birds and rodents.

4. Competitive exclusion may be used in day old chicks to reduce colonisation by Salmonella.

When used, competitive exclusion should be administered according to the instructions provided by the manufacturer and in accordance with the standards and recommendations of the Veterinary Services.

5. Vaccines are used against Salmonella infections caused by different serotypes in various poultry species, including single or combined vaccines. Vaccines produced according to the Terrestrial Manual should be used.

If live vaccines are used it is important that field and vaccine strains be easily differentiated in the laboratory. If serology is used as the surveillance method, it may not be possible to distinguish between vaccination and infection with a field strain.

Vaccination can be used as part of an overall Salmonella control programme. It is recommended that vaccination not be used as the sole control measure.

When the status of the breeding farm and the hatchery from which the flock originates is not known or does not comply with this Chapter, vaccination of flocks starting with day-old chicks, against the Salmonella serotypes known to be significant should be considered.

Vaccination against the Salmonella serotypes known to be significant should be considered when moving day-old chicks to a previously contaminated shed so as to minimise the risk of the birds contracting Salmonella infection.

When used, vaccines should be administered according to the instructions provided by the manufacturer and in accordance with the standards and recommendations of the Veterinary Services.

Vaccination against S. Enteritidis can cause a positive reaction in Salmonella Gallinarum serological tests and needs to be considered when implementing measures for these pathogens.
Annex XIII (contd)

6. Depending on animal health, risk assessment, and public health policies, culling is an option to manage infected breeder and layer flocks. Infected flocks should be destroyed or slaughtered and processed to minimise human exposure to Salmonella.

If poultry are not culled, eggs for human consumption should be diverted for processing for inactivation of Salmonella spp.

7. S. Enteritidis is characterised by its ovarian transmission pattern. Countries should set targets for eradicating (or significantly reducing) Salmonella Enteritidis from egg-producing flocks through a guided policy for eradication from the top of the production pyramid, i.e. from grandparent flocks through breeder flocks to layer flocks.

8. As far as the veterinary involvement is concerned, the responsible veterinarian should monitor the results of surveillance testing for Salmonella. This information should be available to the veterinarian before marketing if a veterinary certificate for flock Salmonella status is required. When required by the Competent Authority, the veterinarian or other authorised person should notify the Competent Authority if the presence of Salmonella of the relevant serotype is confirmed.

Article X.X.6.

Prevention of Salmonella spread from infected flocks

If a flock is found infected with specific Salmonella serotypes of concern, the following actions should be taken in addition to general measures detailed in Chapter 6.3. Hygiene and Biosecurity Procedures in Poultry Production:

1. According to the epidemiological situation, investigations should be carried out to determine the origin of the infection.

2. Movement of poultry flocks at the end of the production cycle should only be allowed for slaughter or destruction. Special precautions should be taken in the transport, slaughter and processing of the birds, e.g. they could be sent to a separate slaughterhouse or processed at the end of a shift before cleaning and disinfection of the equipment.

3. Litter should not be reused. Poultry litter/faeces and other potentially contaminated farm waste should be disposed of in a safe manner to prevent the direct or indirect exposure of humans, livestock and wildlife to Salmonella. Particular care needs to be taken in regard to poultry litter/faeces used to fertilise plants intended for human consumption. If litter is not removed then it should be treated in a manner to inactivate infectious agents, to prevent the spread from one flock to the next.

4. Particular care should be taken in cleaning and disinfection of the poultry house and equipment.

5. Before restocking the facility, a bacteriological examination should be carried out as detailed in this Chapter and the Terrestrial Manual.
CHAPTER 6.X.

INTRODUCTION TO THE RECOMMENDATIONS FOR CONTROLLING ANTIMICROBIAL RESISTANCE

Article 6.X.1.

The purpose of this Chapter is to provide methodologies for OIE Members to appropriately address the emergence or spread of resistant bacteria from the use of antimicrobial agents in animal husbandry and to contain antimicrobial resistance through controlling the use of antimicrobial agents.

Antimicrobial agents are essential drugs 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, controlling and preventing infectious diseases in animals, and this contributes to human health through the supply of animal protein without the risk of transmission of food-borne diseases. The OIE therefore considers that ensuring continued access to effective antimicrobial agents is a priority.

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 prevent or minimise pressures for the selection of antimicrobial resistance factors in humans and animals. Arising from its mandate for the protection of animal health and food safety, the OIE developed these Chapters to provide guidance to Members in regard to risks in the animal sector.

The application of risk management measures should be based on risk assessment that is international standards on microbiological risk analysis and supported by sound data and information when available. The methodologies provided in these Chapters should be consulted as part of the standard approach to preventing and reducing antimicrobial resistance.


**Preamble:** The scope of these recommendations is to deal with stray and feral dogs, which pose serious human health, animal health and welfare problems and have a socio-economic, political, and religious problems in many countries. Whilst acknowledging human health is a priority including the prevention of zoonotic diseases notably rabies, the OIE recognises the importance of controlling dog populations without causing unnecessary or avoidable animal suffering. Veterinary Services should play a lead role in preventing zoonotic diseases and ensuring animal welfare and should be involved in dog population control, coordinating their activities with other competent public institutions and/or agencies.

**Article 7.X.1.**

**Guiding principles**

The following recommendations are based on those laid down in Chapter 7.1. of the OIE Terrestrial Animal Health Code (*The Terrestrial Code*). Some additional principles are relevant to these recommendations:

1. The promotion of responsible dog ownership can significantly reduce the numbers of stray dogs and the incidence of zoonotic diseases.

2. Because dog ecology is linked with human activities, control of dog populations has to be accompanied by changes in human behaviour to be effective.

**Article 7.X.2.**

**Definitions**

**Stray dog**

means any dog not under direct control by a person or not prevented from roaming.

Types of stray dog:

a) free-roaming owned dog not under direct control or restriction at a particular time;

b) free-roaming dog with no owner;

c) feral dog: domestic dog that has reverted to the wild state and is no longer directly dependent upon humans for successful reproduction.

**Owned dog**

means a dog with a person that claims responsibility.

**Person**

this can include more than one individual, and could comprise family/household members or an organisation.
**Responsible dog ownership**

means the situation whereby a person (as defined above) accepts and commits to perform various duties according to the legislation in place and focused on the satisfaction of the behavioral, psychological, environmental and physical needs of a dog and to the prevention of risks (aggression, disease transmission or injuries) that the dog may pose to the community, other animals or the environment.

**Euthanasia**

means the act of inducing death in a humane manner.

**Dog population control programme**

means a programme with the aim of reducing a stray dog population to a particular level and/or maintaining it at that level and/or managing it in order to meet a predetermined objective (see Article 2).

**Carrying capacity**

means is the upper limit of the dog population density that could be supported by the habitat based on the availability of resources (food, water, shelter), and human acceptance.

### Article 7.X.3.

**Dog population control programme objectives**

The objectives of a programme to control the dog population may include the following:

1. improve health and welfare of owned and stray dog population;
2. reduce numbers of stray dogs to an acceptable level;
3. promote responsible ownership;
4. assist in the creation and maintenance of a rabies immune or rabies-free dog population;
5. reduce the risk of zoonotic diseases other than rabies;
6. manage other risks to human health (e.g., parasites);
7. prevent harm to the environment and other animals;
8. prevent illegal trade and trafficking.

### Article 7.X.4.

**Responsibilities and competencies**

1. **Veterinary Authority**

   The Veterinary Authority is responsible for the implementation of animal health and animal welfare legislation, in coordination with other competent government agencies and institutions. Control of endemic zoonotic diseases such as rabies and parasitic infections (e.g., *Echinococcus* spp.) would require technical advice from the Veterinary Authority, as animal health and some aspects of public health are within this Authority’s competence but organising and/or supervising dog control schemes can be the responsibility of non-governmental organisations and governmental agencies other than the Veterinary Authority.
2. Other government agencies

The responsibilities of other government agencies will depend on the risk being managed and the objective/ nature of the dog population control measures employed.

The ministry or other agency responsible for public health would normally play a leadership role and may have legislative authority in dealing with zoonotic diseases. Control of stray dogs with regard to other human health risks (e.g. stray dogs on roads; dog attacks within communities) may fall within the responsibility of the public health agency but is more likely to be the responsibility of the local government authorities of police or other agencies for public safety/security operating at the state/provincial or municipal level.

Environment protection agencies may take responsibility for control problems associated with stray dogs when they present a hazard to the environment (e.g. control of feral dogs in national parks; prevention of dog attacks on wildlife or transmission of diseases to wildlife) or where a lack of environmental controls is giving rise to stray dog populations that threaten human health or access to amenities. For example, environmental protection agencies may regulate and enforce measures to prevent dogs (and other wild animals) from accessing waste or human sewage.

3. Private sector veterinarians

The private sector veterinarian is responsible for providing advice to dog owners or handlers consulting the veterinarian for advice or treatment of a dog. The private sector veterinarian can play an important role in disease surveillance because he/she might be the first to see a dog suffering from a notifiable disease such as rabies. It is necessary that the private sector veterinarian follow the procedure established by the Veterinary Authority for responding to and reporting a suspected rabies case or a dog that is suffering from any other notifiable disease. Private sector veterinarians also play an important role (often in liaison with the police and/or local authorities) in dealing with cases of neglect that can lead to problems with stray and mismanaged dogs.

The private veterinarian has competence and will normally be involved in dog health programmes and population control measures, including health testing, vaccination, identification, kennelling during the absence of the owner, sterilisation and euthanasia. Two-way communication between the private sector veterinarian and Veterinary Authority, often via the medium of a veterinary professional organisation, is very important and the Veterinary Authority is responsible to set up for setting up appropriate mechanisms for this action.

4. Non governmental organisations (NGOs)

Non governmental organisations (NGOs) are potentially important partners of the Veterinary Services in contributing to public awareness and understanding and helping to obtain resources to contribute in a practical way to the design and successful implementation of dog control programmes. NGOs can supply local knowledge on dog populations and features of ownership, as well as expertise in handling and kennelling dogs and the implementation of sterilisation programmes. NGOs can also contribute, together with veterinarians and the authorities in educating the public in responsible dog ownership.

5. Local government authorities

Local government authorities are responsible for many services and programmes that relate to health, safety and public good within their jurisdiction. In many countries the legislative framework gives authority to local government agencies in regard to aspects of public health, environmental health/ hygiene and inspection/ compliance activities.
In many countries local government agencies are responsible for the development and enforcement of legislation relating to dog ownership (e.g., registration, microchipping, vaccination, leash laws, abandonment), the control of stray dogs (e.g., dog catching and shelters) and the alleviation of the problems stray dogs cause in their jurisdiction. This would normally be done with advice from a higher level (national or state/provincial) authority with specialised expertise in regard to public health and animal health. Collaboration with the private sector veterinarians (e.g., in programs to sterilise and vaccinate stray dogs) and NGOs is a common feature of dog control programmes. Regardless of the legislative basis, it is essential to have the co-operation of local government authorities in the control of stray dogs.

6. Dog owners

When a person takes on the ownership of a dog there should be an immediate acceptance of responsibility for that dog, and for any offspring it may produce, for the duration of its life or until a subsequent owner is found. The owner must ensure that the welfare of the dog, including behavioural needs, are respected and the dog is protected, as far as possible, from infectious diseases (e.g., through vaccination and parasite control) and from unwanted reproduction (e.g., through surgical contraception or sterilisation). Owners should ensure that the dog’s ownership is clearly identified (preferably with permanent identification such as a tattoo or microchip) and, where required by legislation, registered on a centralised database. All reasonable steps should be taken to ensure that the dog does not roam out of control in a manner that would pose a problem to the community and/or the environment.

Article 7.X.5.

In the development of a dog population control programme it is recommended that the authorities establish an advisory group, which should include veterinarians, experts in dog ecology, dog behaviour and zoonotic diseases, and representatives of relevant stakeholders (local authorities, human health services/authorities, environmental control services/authorities, NGOs and the public). The main purpose of this advisory group would be to analyse and quantify the problem, identify the causes, obtain public opinion on dogs and propose the most effective approaches to use in the short and long term.

Important considerations are as follows:

1. Identifying the sources of stray dogs
   a) Owned animals dogs that roam freely
   b) Dogs that have been abandoned by their owner, including puppies resulting from uncontrolled breeding of owned dogs.
   c) Unowned dogs that reproduce successfully.

2. Estimating the existing number, distribution and ecology

Practical tools that are available include registers of dogs, population estimates, and surveys of dogs, owners, dog shelters and associated veterinarians. The important factors relevant to the dog carrying capacity of the environment include food, shelter, water and human attitudes and behaviour.

A methodology could be established to make an estimate of the total dog population. An overview of appropriate methodologies may be found in Annex I. The same methodology could be used at appropriate intervals to assess population trends.
3. **Legislation** Regulatory framework

A regulatory framework that would help authorities establish successful dog control programmes could include the following key elements:

a) registration and identification of dogs and licensing of dog breeders;

b) vaccination against rabies and other preventive measures against zoonotic disease, as appropriate;

c) veterinary procedures (e.g. surgical procedures);

d) control of dog movement (national and international);

e) control of dangerous dogs;

f) regulations on the breeding and sale of dogs;

g) environmental controls (e.g. abattoirs, rubbish dumps, dead stock facilities);

h) regulations for dog shelters;

i) animal welfare obligations of owners and authorities.

4. **Resources available to authorities**

a) Human resources;

b) financial resources;

c) technical tools;

d) infrastructure;

e) cooperative activities;

f) public-private-NGO partnerships;

g) central-state or province-local partnerships.

**Article 7.X.6.**

**Control measures**

The following control measures could be implemented according to the national context and local circumstances. Measures may be used in combination. Euthanasia of dogs, used alone, is not an effective control measure. If used, it should be done humanely (see Article 5.11) and in combination with other measures to achieve effective long term control. It is also important that authorities gain an understanding of people’s attitudes towards dog ownership so that they can develop a cooperative approach to the control of dog populations.

1. **Education and legislation for responsible ownership**

Encouraging dog owners to be more responsible will reduce the number of dogs allowed to roam, improve the health and welfare of dogs, and minimise the risk that dogs pose to the community. The promotion of responsible dog ownership through legislation and education is a necessary part of a dog population control programme. Collaboration with local government authorities, animal welfare NGOs, kennel clubs, private veterinarians and veterinary organisations will assist Veterinary Authorities in establishing and maintaining programmes.
Annex XV (contd)

Education on responsible dog ownership (for the currently owned dog and any offspring it produces) should address the following elements:

a) the importance of proper selection and care to ensure the welfare of the dog and any offspring; the latter may include preparing the dog to cope with its environment through attention to socialisation and training;

b) registration and identification of dogs (see Article 5.2.);

c) disease prevention, in particular zoonotic disease, e.g. through regular vaccination in rabies endemic areas;

d) preventing negative impacts of dogs on the community, via pollution (e.g. faeces and noise), risks to human health through biting or traffic accidents and risks to other dogs, wildlife, livestock and other companion animal species;

e) control of dog reproduction.

In order to achieve a shift towards responsible ownership, a combination of legislation, public awareness, education, and promotion of these elements will be required. It may also be necessary to improve access to resources supporting responsible ownership, such as veterinary care, identification and registration services and measures for control of zoonotic diseases.

2. Registration and identification of dogs (licensing)

A core component of dog population control by the Competent Authorities is the registration and identification of owned dogs. This may include granting licences to owners and breeders. Registration and identification may be emphasized as part of responsible dog ownership and are often linked to animal health programs, for example, mandatory rabies vaccination and a dog traceability.

Registration of animals in a centralised database can be used to support the enforcement of legislation and the reuniting of lost animals with owners. The control of dog reproduction by sterilisation can be encouraged through financial incentives presented by differential licensing fees.

3. Reproductive control

Controlling reproduction in dogs prevents the birth of unwanted puppies and can help address the balance between demand for dogs and the size of the population. It is advisable to focus efforts to control reproduction on those individuals or groups in the dog population identified as the most productive and the most likely to be the sources of unwanted and stray dogs, to ensure best use of resources. Methods of controlling reproduction will require direct veterinary input to individual animals. Involvement of both private and public veterinary sectors may be required to meet demand for services. Subsidisation of sterilisation programmes by government or other organisations may be considered to encourage uptake. The control of reproduction is essentially the responsibility of owners and can be incorporated into education on responsible ownership (section 5a.). Methods for controlling reproduction in dogs include:

a) surgical sterilisation;

b) chemical sterilisation;

c) chemical contraception;

d) separation of female dogs during oestrus from unsterilised males.
Surgical sterilisation should be carried out by a veterinarian and include appropriate anaesthesia and pain management relief.

Any chemicals or drugs used in controlling reproduction should be shown to have appropriate safety, quality and efficacy for the function required and used according to the manufacturer’s and Competent Authority’s regulations. In the case of chemical sterilants and contraceptives, research and field trials may need to be completed before use.

4. Removal and handling

The Competent Authority should collect dogs that are not under direct supervision and verify their ownership. Capture, transport, and holding of the animals should be done humanely. The Competent Authority should develop and implement appropriate legislation and training to regulate these activities. Capture should be achieved with the minimum force required and equipment should be used that supports humane handling. Uncovered wire loops should not be used for capture.

5. Management of captured stray dogs: Capture and return, rehoming or release

Competent Authorities have the responsibility to develop minimum standards for the housing (physical facilities) and care of these dogs. There should be provision for holding the dogs for a reasonable period of time to allow for reunion with the owner and, as appropriate, for rabies observation.

a) Minimum standards for housing should include the following provisions:
   i) site selection: Access to drainage, water and electricity are essential and environmental factors such as noise and pollution should be taken into account;
   ii) kennel size, design and occupancy taking exercise into account;
   iii) disease control measures including isolation and quarantine facilities.

b) Management should address:
   i) adequate fresh water and nutritious food;
   ii) regular hygiene and cleaning;
   iii) routine inspection of the dogs;
   iv) monitoring of health and provision of required veterinary treatments;
   v) policies and procedures for rehoming (adoption), sterilisation and euthanasia;
   vi) training of staff in safe and appropriate handling of dogs;
   vii) record keeping and reporting to authorities.

Dogs that are removed from a community may be reunited with the owner or offered to new owners for adoption (rehoming). This provides an opportunity to promote responsible ownership and good animal health care (including rabies vaccination). Prior to adoption, rehoming, authorities may consider sterilisation of dogs as a population control measure. Dogs should be sterilised. The suitability of new owners to adopt dogs should be assessed and owners matched with available animals. The effectiveness of rehoming may be limited due to the suitability and number of dogs.
Dogs that are removed from a community may in some cases be provided with health care (including rabies vaccination), sterilised, and released to their local community at or near the place of capture. This method is more likely to be accepted in the situation where the presence of stray dogs is considered to be inevitable and is well tolerated by the local community.

This method is not applicable in all situations and may be illegal in countries or regions where legislation prohibits the abandonment of dogs. Problems caused by dogs, such as noise, faecal pollution, bite injuries and traffic accidents, would not be alleviated as dogs are returned to the local community and their movements are not restricted. If the local community has owned dogs, and sterilised dogs are released, consideration should be given to the risk that this could encourage abandonment of unwanted dogs. In the situation where many dogs are owned, a population control programme that focuses on neutering and responsible ownership may be more appropriate.

It is recommended that before adopting this approach, a cost-benefit analysis is conducted. Factors such as the monetary costs, impact on culture of ownership and public safety should be assessed as well as the benefits for disease control and animal welfare as well as any societal benefits.

c) If this method is adopted, the following factors should be addressed:

i) raising awareness of the programme within the local community to ensure understanding and support;

ii) use of humane methods for catching, transporting and holding dogs;

iii) correct surgical technique, anaesthesia and analgesia, followed by post-operative care;

iv) disease control may include blanket vaccination (e.g. rabies) and treatments and testing for diseases (e.g. leishmaniasis) followed, as appropriate by treatment or euthanasia of the dog;

v) behavioural observation may be used to assess if dogs are suitable for release; if not suitable for release or rehoming, euthanasia should be considered;

vi) permanent marking (e.g. tattoo or microchip) to indicate that the animal has been sterilised. Individual identification also allows for tracking of vaccination status and treatment history and identification of a level of ‘ownership’ by the organisation/authority responsible for carrying out this intervention; a visible identification (e.g. collar) may also be used to prevent unnecessary recapture; identification can also be taken to indicate a level of ‘ownership’ by the organisation/authority responsible for carrying out this intervention;

vii) the dog should be returned to a place that is as near as possible to the place of capture;

viii) the welfare of dogs after release should be monitored and action taken if required.

Dogs that are removed from a community may be too numerous or may be unsuitable for any rehoming scheme. If euthanasia of these unwanted animals is the only option, the procedure should be conducted in accordance with the regulations of the Competent Authority (see Article 5.11).

5. Environmental controls

Steps should be taken to reduce the carrying capacity, such as excluding dogs from sources of food (e.g. rubbish dumps and abattoirs, and installing animal-proof rubbish containers).
This should be linked to a reduction in the animal dog population by other methods, to avoid animal welfare problems.

7. Control of dog movement – international (export/ import)

Chapter 8.11.2.5 of the Terrestrial Animal Health Code provides recommendations on the international movement of dogs between rabies free countries and countries considered to be infected with rabies.

8. Control of dog movements – within country (e.g. leash laws, roaming restrictions)

Measures for the control of dog movement in a country are generally invoked for the following reasons:

a) for rabies control when the disease is present in a country;

b) for public safety reasons;

c) for the safety of “owned dogs” in an area or locality when a stray dog control programme is in place;

d) to protect wildlife and livestock.

It is necessary to have a empowering regulatory framework legislation to give the necessary power is necessary and a national or local infrastructure comprising organization, administration, staff and resources to encourage the finders of a stray dog to report to the Competent Authority.

9. Regulation of commercial dog dealers

Dog breeders and dealers should be encouraged to form or join an appropriate association. Such associations should encourage a commitment to the raising and selling of physically and psychologically healthy dogs, as unhealthy dogs may be more likely to be abandoned to become part of the stray population. They should encourage breeders and dealers to provide advice on proper care to all new owners of dogs. Regulations covering commercial dog breeders and dealers should include specific requirements for accommodation, provision of suitable food, drink and bedding, adequate exercise, veterinary care and disease control and may require breeders and dealers to allow regular inspection, including veterinary inspection.

10. Reduction in dog bite incidence

The most effective means of reducing prevalence of dog bites are education and placing responsibility on the owner. Dog owners should be educated in principles of responsible dog ownership as described in Article 5.1. Legal mechanisms that enable the Competent Authorities to impose penalties or otherwise deal with irresponsible owners are necessary. Mandatory registration and identification schemes will facilitate the effective application of such mechanisms. Young children are the group at highest risk for dog bites. Public Education programmes focussed on appropriate dog-directed behaviour have been demonstrated to be effective in reducing dog bite prevalence and these programmes should be encouraged. Authorities should seek advice from dog behaviour experts in developing dog safety education programmes.
11. **Euthanasia**

When euthanasia is practised, the general principles in the Code should be followed, with the emphasis on using the most practical, rapid and humane methods and ensuring operator safety. For practical reasons, different procedures may be used in rural and urban areas. Regardless of the method used, it is important to minimise distress, anxiety and pain by ensuring that operators are appropriately trained.

Table 1 shows a list of methods for the euthanasia of dogs.
### Table 1: List of methods for the euthanasia of dogs

<table>
<thead>
<tr>
<th>Euthanasia method</th>
<th>Specific method</th>
<th>Animal welfare concerns/implications</th>
<th>Key animal welfare requirements</th>
<th>Considerations relating to operator security</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chemical</strong></td>
<td>Barbiturates</td>
<td>Correct restraint is needed.</td>
<td>Recommend to use IV injection.</td>
<td>Correct restraint is needed.</td>
<td>Speed of action generally depends on the dose, concentration, route and rate of injection.</td>
<td>These drugs persist in the carcass and may cause sedation or death in animals that consume the cadaver.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>IP is slow and may be irritant.</td>
<td>When using IP injection, the solution may be diluted or local anaesthetic agent used in conjunction.</td>
<td>Administered under veterinary supervision and requires trained personnel.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>IC injection is a painful procedure.</td>
<td>IC should only be performed on unconscious animal and by skilled operator.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Embutramide + Mebezonium + Tetracaine</td>
<td>Muscle paralysis may occur before loss of consciousness if injection given rapidly</td>
<td>Use slow IV injection with sedation to permit slow rate of injection.</td>
<td>Correct restraint is needed.</td>
<td>Quite low cost.</td>
<td>Unavailable/unlicensed in some countries</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>To be administered under veterinary supervision and by trained personnel.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anaesthetic agent overdose (thiopentone or propofenol)</td>
<td>Underdosing may lead to recovery</td>
<td>IV injection of a sufficient dose</td>
<td>Correct restraint is needed.</td>
<td>Generally quick action and minimal discomfort to animal.</td>
<td>Large volume required (cost implications)</td>
</tr>
<tr>
<td></td>
<td>Potassium chloride (K Cl)</td>
<td>K+ is cardiotoxic and very painful if used without anaesthetic agent.</td>
<td>Only use on anaesthetised animals, IV injection</td>
<td>Requires trained personnel.</td>
<td>Readily available without veterinary control.</td>
<td>Prior need for anaesthetic (cost and availability implications)</td>
</tr>
</tbody>
</table>
### Table 1: List of methods for the euthanasia of dogs (contd)

<table>
<thead>
<tr>
<th>Method</th>
<th>Considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Free bullet</strong></td>
<td>Can be inhumane if shot is inaccurate and dog is only wounded; dog may also escape.</td>
</tr>
<tr>
<td>Skilled operator essential.</td>
<td>Risk of injury to operators and spectators.</td>
</tr>
<tr>
<td>Not necessary to handle or capture dog.</td>
<td>Brain tissue may be unavailable for rabies diagnosis. Risk of injury to bystanders. Legal constraints on use of firearms.</td>
</tr>
<tr>
<td><strong>Penetrating captive bolt</strong></td>
<td>Can be inhumane if shot is inaccurate and dog is only wounded.</td>
</tr>
<tr>
<td>Skilled operator essential.</td>
<td>Animal must be restrained. Skilled operator essential.</td>
</tr>
<tr>
<td>No risk to operator (cf free bullet) unless risk of dog infected with rabies, due to potential contact with brain tissue</td>
<td>Brain tissue may be unavailable for rabies diagnosis. Legal constraints on use of firearms. May raise aesthetic objections.</td>
</tr>
<tr>
<td><strong>Exsanguination</strong></td>
<td>Onset of hypovolaemia may cause dog to become anxious.</td>
</tr>
<tr>
<td>Only use on unconscious animal</td>
<td>Danger to operator through use of sharp instrument.</td>
</tr>
<tr>
<td>Material requirements minimal.</td>
<td>Must be done on unconscious animal. Aesthetically objectionable.</td>
</tr>
<tr>
<td>Aesthetically objectionable.</td>
<td></td>
</tr>
</tbody>
</table>

**Mechanical**
Table 1: List of methods for the euthanasia of dogs (contd)

<table>
<thead>
<tr>
<th>Euthanasia method</th>
<th>Specific method</th>
<th>Animal welfare concerns/implications</th>
<th>Key animal welfare requirements</th>
<th>Considerations relating to operator security</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gaseous</td>
<td>Carbon monoxide (CO)</td>
<td>Inadequate concentration of CO is not lethal and can cause suffering. Signs of distress (convulsions, vocalization and agitation) may occur.</td>
<td>Compressed CO in cylinders must be used to achieve and maintain adequate concentration, which must be monitored. Note: fumes from gasoline engines are an irritant and this source of CO is not recommended.</td>
<td>Very hazardous for operator - gas is odourless and causes toxicity at both acute high levels and chronic low levels</td>
<td>Dog dies quite rapidly if concentration of 4 to 6% used. No odour (therefore no aversive effect). Gas is not flammable or explosive except at concentration greater than 10%.</td>
<td></td>
</tr>
</tbody>
</table>
### Table 1: List of methods for the euthanasia of dogs (contd)

<table>
<thead>
<tr>
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<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gaseous</strong></td>
<td>Carbon dioxide (CO&lt;sub&gt;2&lt;/sub&gt;)</td>
<td>Gas is aversive. Inadequate concentration of CO&lt;sub&gt;2&lt;/sub&gt; is not lethal and can cause suffering. CO&lt;sub&gt;2&lt;/sub&gt; is heavier than air, so when incomplete filling of the chamber occurs, dogs may raise their head and avoid exposure. Few studies on adequate concentration and animal welfare.</td>
<td>Compressed CO&lt;sub&gt;2&lt;/sub&gt; gas chamber is the only acceptable recommended method because the concentration can be monitored and regulated.</td>
<td>Minimal hazard to operator when properly designed equipment used.</td>
<td>Gas is not flammable or explosive and causes quite rapid anaesthesia when correct concentrations used. Low cost. Readily available as compressed gas.</td>
<td>Unconsciousness can occur in minutes, but death may take some time. Likelihood of suffering before unconsciousness.</td>
</tr>
<tr>
<td></td>
<td>Inert gas (nitrogen, N&lt;sub&gt;2&lt;/sub&gt;, argon, Ar)</td>
<td>Loss of consciousness is preceded by hypoxemia and ventilatory stimulation, which may be distressing to the dog. Re-establishing a low concentration of O&lt;sub&gt;2&lt;/sub&gt; (i.e. greater than or equal to 6%) in the chamber before death will allow immediate recovery.</td>
<td>Concentration above 98% must be achieved rapidly and maintained. Properly designed equipment must be used</td>
<td>Minimal hazard to operator when properly designed equipment used.</td>
<td>Gas is not flammable or explosive and is odourless. Readily available as compressed gas.</td>
<td>High cost. Little data on animal welfare implications in dogs.</td>
</tr>
</tbody>
</table>
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</tr>
</thead>
<tbody>
<tr>
<td>Gaseous</td>
<td>Anaesthetic gas overdose (halothane or enflurane)</td>
<td>Animal may struggle and become anxious during induction. Vapours may be irritating and can induce excitement.</td>
<td>Supplementation with air or O₂ required to avoid hypoxemia during induction phase.</td>
<td>Some gases may be hazardous, especially for pregnant women. General recommendation: Avoid human exposure to greater than or equal to 2ppm to avoid narcosis.</td>
<td>Gas is not flammable or explosive. Valuable for use with small animals (&lt;7kgs) and animals that are already anesthetised with gas.</td>
<td>High cost. Anaesthetic and euthanasia properties of the gas used must be known. Isoflurane has a pungent odour. Methoxyflurane's action is slow and dog may become agitated.</td>
</tr>
<tr>
<td>Electrical</td>
<td>Electrocution</td>
<td>Cardiac fibrillation occurs before onset of unconsciousness, causing severe pain if dog is conscious. Pain can also be caused by violent extension of the limbs, head and neck. Method may not be effective if insufficient current applied.</td>
<td>Dogs must be unconscious before being electrocuted. This can be accomplished by electrical stunning (current through the brain to produce an instantaneous stun) or anaesthesia. Electrodes should span the brain in order that the current passed through the brain in order to achieve an effective stun. Death would result from current passed through the heart of an unconscious animal. Proper equipment and trained operator is essential.</td>
<td>May be hazardous for operator, who should use protective equipment (boots and gloves).</td>
<td>Low cost.</td>
<td>Inhumane if performed on conscious dog. May raise aesthetic objections.</td>
</tr>
</tbody>
</table>

KEY to abbreviations used in Table 1:
IV: intravenous
IP: Intraperitoneal
IC: Intracardiac
Annex XV (contd)

a) Comments on methods for the euthanasia of dogs:

i) Restraint

When a dog needs to be restrained for any procedure, including euthanasia, this should always be done with full regard for operator security and animal welfare. Some euthanasia methods must be used in association with sedation or anaesthesia in order to be considered humane.

ii) Special equipment

When special equipment is needed to perform euthanasia (e.g. gas chamber) the system should be designed for the purpose and regularly maintained in order to achieve operator security and animal welfare.

iii) The following methods, procedures and practices are unacceptable on animal welfare grounds:

- **Chemical methods:**
  - Embutramide + Mebezonium + Tetracaine without sedation or by other than IV injection
  - Chloral hydrate
  - Nitrous oxide: may be used with other inhalants to speed the onset of anaesthesia, but alone it does not induce anaesthesia in dogs
  - Ether
  - Chloroform
  - Cyanide
  - Strychnine
  - Neuromuscular blocking agents (nicotine, magnesium sulphate, potassium chloride, all curariform agents): when used alone, respiratory arrest occurs before loss of consciousness, so the dog may perceive pain
  - Formalin
  - Household products and solvents.

- **Mechanical methods:**
  - Air embolism on conscious animal
  - Burning
  - Exsanguination of conscious animal
  - Decompression: expansion of gas trapped in body cavities may be very painful
  - Drowning
  - Hypothermia, rapid freezing
  - Stunning: stunning is not a euthanasia method, it should always be followed by a method which ensures death.
  - Kill-trapping
  - Electrocution of conscious animal.
Because neonatal animals and adults with impaired breathing or low blood pressure are resistant to hypoxia, methods that depend upon achieving a hypoxic state (e.g. CO₂, CO, N₂, Ar) should not be used. These methods should not be used in animals aged less than 2 months, except to produce loss of consciousness and should be followed by another method to cause death. Concussion and cervical dislocation may be used in very small neonatal dogs and only in cases of emergency.

Operators must be well trained in the use of physical techniques to ensure that they are correctly and humanely carried out. The dog must be exsanguinated immediately after concussion or cervical dislocation.

iv) Confirmation of death

For all methods of euthanasia used, death must be confirmed before animals are disposed of or left unattended. If an animal is not dead, another method of euthanasia must be performed.

v) Carcass disposal

Carcasses should be disposed of in a manner that complies with legislation. Attention must be paid to the risk of residues occurring in the carcase. Incineration is generally the safest way of carcass disposal.

Article 7.X.7

Monitoring and evaluation of dog population control programmes

Monitoring and evaluation allows for comparison of important indicators against the baselines measured during initial assessment (Article 4). The three main reasons for carrying out monitoring and evaluation are:

1. to help improve performance, by highlighting both problems and successful elements of interventions;

2. for accountability, to demonstrate that the programme is achieving its aims;

3. assuming methods are standardised, to compare the success of strategies used in different locations and situations.

Monitoring is a continuous process that aims to check the programme progress against targets and allows for regular adjustments. Evaluation is a periodic assessment, usually carried out at particular milestones to check the programme is having the desired and stated impact. These procedures involve the measurement of 'indicators' that are chosen because they reflect important components of the programme at different stages. Selection of suitable indicators requires clear planning of what the programme is aiming to achieve, the best selection of indicators will be one that reflects the interest of all relevant stakeholders. Standardised methodology will facilitate comparison of data from subsequent evaluations and performance between different projects. Indicators can be direct measurements of an area targeted to change (e.g. population of free roaming dogs on public property) or indirect measures that reflect change in a targeted area (e.g. number of reported dog bites as a reflection of rabies prevalence).
4. Elements that should generally be monitored and evaluated include:

a) dog population size, separated into sub-populations according to ownership and restriction of movement (i.e. roaming unrestricted or restricted by an owner);

b) dog welfare, in the target population (e.g. body condition score, skin conditions and injuries or lameness) and as a result of the programme (if interventions involve direct handling of dogs, the welfare of the dogs as result of this handling should be monitored);

c) prevalence of zoonotic diseases, such as rabies, in both the animal and human population;

d) responsible animal ownership, including measures of attitudes and understanding of responsible ownership and evidence that this is translating into responsible behaviour.

5. There are many sources of information for measuring indicators monitoring and evaluation purposes, including:

a) feedback from the local community (e.g. through the use of structured questionnaires, focus groups or ‘open format’ consultation processes);

b) records and opinions obtained from relevant professionals (e.g. veterinarians, medical doctors, law enforcement agencies, educators);

c) animal based measurements (e.g. direct observation surveys of population size and welfare status).

The output of activities against budget should be carefully recorded in order to evaluate the effort (or cost) against the outcomes and impact (or benefit) that are reflected in the results of monitoring and evaluation.
Annex XV (contd)

Annex I:

An overview of appropriate methodologies for estimating the size of dog populations.

Population estimates are necessary for making realistic plans for dog population management and zoonosis control, and for monitoring the success of such interventions. However, for designing effective management plans, data on population sizes alone are insufficient. Additional information is required, such as degrees of supervision of owned dogs, the origin of ownerless dogs, accessibility, etc.

The term “owned” may be restricted to a dog that is registered with licensing authorities, or it may be expanded to unregistered animals that are somewhat supervised and receive shelter and some form of care in individual households. Owned dogs may be well supervised and restrained at all times, or they may be left without control for various time periods and activities. Dog owners that claim responsibility may still be accepted or tolerated in the neighbourhood, and individuals may provide food and protection. Such animals are sometimes called “community owned dogs” or “neighbourhood dogs”. For an observer it is frequently impossible to decide if a free roaming dog belongs to someone or not.

The choice of methods for assessing the size of a dog population depends on the ratio of owned versus ownerless dogs, which may not always easy to judge. For populations with a large proportion of owned dogs it may be sufficient to consult dog registration records or to conduct household surveys. These surveys should establish the number of owned dogs and the dog to human ratio in the area. In addition, questions on dog reproduction and demographics, care provided, zoonosis prevention, dog bite incidence, etc. may be asked. Sample questionnaires can be found in the “Guidelines for Dog Population Management” (WHO/WSPA 1990). Standard polling principles must be applied.

If the proportion of ownerless dogs is high or difficult to assess, then one must resort to more experimental approaches. Methods borrowed from wildlife biology can be applied. These methods are described WHO/WSPA’s “Guidelines for Dog Population Management” (1990), and in more detail in numerous professional publications and handbooks, such as Bookhout (1994) and Sutherland (2006). Being generally diurnal and tolerant to human proximity, dogs lend themselves to direct observation and the application of mark-recapture techniques. Nevertheless, a number of caveats and limitations have to be taken into account. Firstly, the risk of zoonotic disease transmission is increased through close physical contact. Also, the methods are relatively labour intensive, they require some understanding of statistics and population biology, and most importantly, they are difficult to apply to very large areas. One must take into account that dog distribution is non-random, that their populations are not static, and that individual dogs are fairly mobile.

Counting of dogs visible in a defined area is the simplest approach to getting information on population size. One has to take into account that the visibility of dogs depends on the physical environment, but also on dog and human activity patterns. The visibility of animals changes with the time of the day and with seasons as a function of food availability, shelter (shade), disturbance, etc. Repeated standardized counting of dogs visible within defined geographical localities (e.g. wards) and specific times will provide indications of population trends. Direct counting is most reliable if it is applied to small and relatively confined dog populations, e.g. in villages, where it might be possible to recognize individual dogs based on their physical appearance.

Methods using mark-recapture procedures are often considered more reliable. However, they also produce trustworthy results only when a number of preconditions are met. Mortality, emigration and recruitment into the population must be minimal during the census period. One may be able to incorporate corrective factors into the calculations.
Annex XV (contd)

It is therefore important that the recommended census procedures are applied at times of low dispersal and that one selects study plots of shape and size that minimize the effect of dog movements in and out of the observation area. Census surveys should be completed within a few days to a maximum of two weeks in order to reduce demographic changes. In addition, all individuals in the population must have an equal chance of being counted. This is a highly improbable condition for dogs, whose visibility depends on ownership status and degrees of supervision. It is therefore recommended that the investigator determines what fraction of the total population he/she might cover with an observational method and how much this part overlaps with the owned dog segment that he/she assesses with household surveys.

There are essentially two ways to obtain a population estimate if it is possible, in a defined area and within a few days, to tag a large number of dogs with a visible mark, e.g. a distinctive collar or a paint smudge. The first method requires that the capture (marking) effort remains reasonably constant for the whole length of the study. By plotting the daily number of dogs marked against the accumulated total of marked dogs for each day one can extrapolate the value representing the total number of dogs in the area. More commonly used in wildlife studies are mark recapture methods (Peterson-Jackson, Lincoln indices). Dogs are marked (tagged) and released back into the population. The population is subsequently sampled by direct observation. The number of marked and unmarked dogs is recorded. One multiplies the number of dogs that were initially marked and released by the number of subsequently observed dogs divided by the number of dogs seen as marked during the re-observation to obtain a total population estimate. Examples for the two methods are given in WHO/WSPA’s “Guidelines for Dog Population Management” (1990).

Since the dog populations of entire countries, states, provinces or even cities are much too large for complete assessment, it is necessary to apply the methods summarized above to sample areas. These should be selected (using common sense) so that results can be extrapolated to larger areas.


CHAPTER 8.5.

FOOT AND MOUTH DISEASE

Article 8.5.1.

Introduction

For the purposes of the Terrestrial Code, the incubation period for foot and mouth disease (FMD) shall be 14 days.

For the purposes of this Chapter, ruminants include animals of the family of Camelidae (except Camelus dromedarius).

For the purposes of this Chapter, a case includes an animal infected with FMD virus (FMDV).

For the purposes of international trade, this Chapter deals not only with the occurrence of clinical signs caused by FMDV, but also with the presence of infection with FMDV in the absence of clinical signs.

The following defines the occurrence of FMDV infection:
1. FMDV has been isolated and identified as such from an animal or a product derived from that animal; or
2. viral antigen or viral ribonucleic acid (RNA) specific to one or more of the serotypes of FMDV has been identified in samples from one or more animals, whether showing clinical signs consistent with FMD or not, or epidemiologically linked to a confirmed or suspected outbreak of FMD, or giving cause for suspicion of previous association or contact with FMDV; or
3. antibodies to structural or nonstructural proteins of FMDV that are not a consequence of vaccination, have been identified in one or more animals showing clinical signs consistent with FMD, or epidemiologically linked to a confirmed or suspected outbreak of FMD, or giving cause for suspicion of previous association or contact with FMDV.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 8.5.2.

FMD free country where vaccination is not practised

Susceptible animals in the FMD free country where vaccination is not practised should be separated from neighbouring infected countries by a buffer zone or physical or geographical barriers, and the application of animal health measures that effectively prevent the entry of the virus, taking into consideration physical or geographical barriers. These measures may include a protection zone should be implemented.

To qualify for inclusion in the existing list of FMD free countries where vaccination is not practised, a Member should:
1. have a record of regular and prompt animal disease reporting;
2. send a declaration to the OIE stating that:
   a) there has been no outbreak of FMD during the past 12 months;
   b) no evidence of FMDV infection has been found during the past 12 months;
Annex XVI (contd)

c) no vaccination against FMD has been carried out during the past 12 months;

d) no vaccinated animal has been introduced since the cessation of vaccination;

3. supply documented evidence that:

a) surveillance for both FMD and FMDV infection in accordance with Articles 8.5.40. to 8.5.46. is in operation;

b) regulatory measures for the early detection, prevention and control of FMD have been implemented.

The Member will be included in the list only after the submitted evidence has been accepted by the OIE. Retention on the list requires that the information in points 2 and 3b) above be re-submitted annually and changes in the epidemiological situation or other significant events should be reported to the OIE according to the requirements in Chapter 1.1.

Article 8.5.3.

FMD free country where vaccination is practised

Susceptible animals in the FMD free country where vaccination is practised should be separated from neighbouring infected countries by a buffer zone or physical or geographical barriers, and the application of animal health measures that effectively prevent the entry of the virus, taking into consideration physical or geographical barriers. These measures may include a protection zone should be implemented.

To qualify for inclusion in the list of FMD free countries where vaccination is practised, a Member should:

1. have a record of regular and prompt animal disease reporting;

2. send a declaration to the OIE that there has been no outbreak of FMD for the past 2 years and no evidence of FMDV circulation for the past 12 months, with documented evidence that:

a) surveillance for FMD and FMDV circulation in accordance with Articles 8.5.40. to 8.5.46. is in operation, and that regulatory measures for the prevention and control of FMD have been implemented;

b) routine vaccination is carried out for the purpose of the prevention of FMD;

c) the vaccine used complies with the standards described in the Terrestrial Manual.

The Member will be included in the list only after the submitted evidence has been accepted by the OIE. Retention on the list requires that the information in point 2 above be re-submitted annually and changes in the epidemiological situation or other significant events should be reported to the OIE according to the requirements in Chapter 1.1.

If a Member that meets the requirements of a FMD free country where vaccination is practised wishes to change its status to FMD free country where vaccination is not practised, the status of this country remains unchanged for a period of at least 12 months after vaccination has ceased. Evidence should also be provided showing that FMDV infection has not occurred during that period.
Article 8.5.4.

**FMD free zone where vaccination is not practised**

A FMD free zone where vaccination is not practised can be established in either a FMD free country where vaccination is practised or in a country of which parts are infected. In defining such zones the principles of Chapter 4.3. should be followed. Susceptible animals in the FMD free zone can should be separated protected from the rest of the country and from neighbouring countries by a buffer zone or by physical/ geographical barriers from the rest of the country and from neighbouring countries if they are of a different animal health status, and by the application of animal health measures that effectively prevent the entry of the virus, taking into consideration physical or geographical barriers. These measures may include a protection zone should be implemented.

A Member in which a FMD free zone where vaccination is not practised is to be established should To qualify for inclusion in the list of FMD free zones where vaccination is not practised, a Member should:

1. have a record of regular and prompt animal disease reporting;

2. send a declaration to the OIE stating that it wishes to establish a FMD free zone where vaccination is not practised, and that within the proposed FMD free zone:
   a) there has been no outbreak of FMD during the past 12 months;
   b) no evidence of FMDV infection has been found during the past 12 months;
   c) no vaccination against FMD has been carried out during the past 12 months;
   d) no vaccinated animal has been introduced into the zone since the cessation of vaccination, except in accordance with Article 8.5.9.;
   e) documented evidence shows that surveillance in accordance with Articles 8.5.40. to 8.5.46. is in operation for both FMD and FMDV infection;

3. describe in detail:
   a) regulatory measures for the prevention and control of both FMD and FMDV infection,
   b) the boundaries of the proposed FMD free zone and, if applicable, the buffer protection zone or physical or geographical barriers,
   c) the system for preventing the entry of the virus (including the control of the movement of susceptible animals) into the proposed FMDV free zone (in particular if the procedure described in Article 8.5.9. is implemented),

and supply documented evidence that these are properly implemented and supervised.

The proposed free zone will be included in the list of FMD free zones where vaccination is not practised only after the submitted evidence has been accepted by the OIE.

The information required in points 2 and 3c) above should be re-submitted annually and changes in the epidemiological situation or other significant events including those relevant to points 3a) and 3b) should be reported to the OIE according to the requirements in Chapter 1.1.
Article 8.5.5.

**FMD free zone where vaccination is practised**

A FMD free zone where vaccination is practised can be established in either a FMD free country where vaccination is not practised or in a country of which parts are infected. In defining such zones the principles of Chapter 4.3. should be followed. Susceptible animals in the FMD free zone where vaccination is practised can be separated from neighbouring countries or zones if they are infected of a lesser animal health status by a buffer zone or by physical/geographical barriers from the rest of the country and from neighbouring countries if they are of a different animal health status, and the application of animal health measures that effectively prevent the entry of the virus, taking into consideration physical or geographical barriers. These measures may include a protection zone should be implemented.

A Member in which a FMD free zone where vaccination is practised is to be established should To qualify for inclusion in the list of FMD free zones where vaccination is practised, a Member should:

1. have a record of regular and prompt animal disease reporting;

2. send a declaration to the OIE that it wishes to establish a FMD free zone where vaccination is practised and that within the proposed FMD free zone:
   a) there has been no outbreak of FMD for the past 2 years;
   b) no evidence of FMDV circulation for the past 12 months;
   c) documented evidence shows that surveillance in accordance with Articles 8.5.40. to 8.5.46. is in operation for FMD and FMDV circulation;

3. supply documented evidence that the vaccine used complies with the standards described in the Terrestrial Manual;

4. describe in detail:
   a) regulatory measures for the prevention and control of both FMD and FMDV circulation,
   b) the boundaries of the proposed FMD free zone where vaccination is practised and, if applicable, the buffer protection zone or physical or geographical barriers,
   c) the system for preventing the entry of the virus into the proposed FMD free zone (in particular if the procedure described in Article 8.5.9. is implemented),

and supply evidence that these are properly implemented and supervised.

The proposed free zone will be included in the list of FMD free zones where vaccination is practised only after the submitted evidence has been accepted by the OIE. The information required in points 2, 3 and 4c) above should be re-submitted annually and changes in the epidemiological situation or other significant events including those relevant to points 4a) and 4b) should be reported to the OIE according to the requirements in Chapter 1.1.
If a Member that has a zone which meets the requirements of a FMD free zone where vaccination is practised wishes to change the status of the zone to FMD free zone where vaccination is not practised, the status of this zone remains unchanged for a period of at least 12 months after vaccination has ceased. Evidence should also be provided showing that FMDV infection has not occurred in the said zone during that period.

Article 8.5.6.

FMD infected country or zone

A FMD infected country is a country that does not fulfil the requirements to qualify as either a FMD free country where vaccination is not practised or a FMD free country where vaccination is practised.

A FMD infected zone is a zone that does not fulfil the requirements to qualify as either a FMD free zone where vaccination is not practised or a FMD free zone where vaccination is practised.

Article 8.5.7.

Establishment of a containment zone within a FMD free country or zone

In the event of limited outbreaks within a FMD free country or zone, including within a protection zone, with or without vaccination, a single containment zone, which includes all cases, can be established for the purpose of minimizing the impact on the entire country or zone.

For this to be achieved, the Veterinary Authority should provide documented evidence that:

1. the outbreaks are limited based on the following factors:
   a) immediately on suspicion, a rapid response including notification has been made;
   b) standstill of animal movements has been imposed, and effective controls on the movement of other commodities mentioned in this Chapter are in place;
   c) epidemiological investigation (trace-back, trace-forward) has been completed;
   d) the infection has been confirmed;
   e) the primary outbreak and likely source of the outbreak has been identified;
   f) all cases have been shown to be epidemiologically linked;
   g) no new cases have been found in the containment zone within a minimum of two incubation periods as defined in Article 8.5.1. after the stamping-out of the last detected case is completed;

2. a stamping-out policy has been applied;

3. the susceptible animal population within the containment zones should be clearly identifiable as belonging to the containment zone;

4. increased passive and targeted surveillance in accordance with Articles 8.5.40. to 8.5.46. in the rest of the country or zone has been carried out and has not detected any evidence of infection;
5. animal health measures that effectively to prevent the spread of the FMDV infection from the containment zone to the rest of the country or zone, including taking into consideration physical and geographical barriers, are in place;

6. ongoing surveillance in the containment zone are in place;

6. containment zone should be large enough to contain the disease and comprise include both a restricted/protection zone and larger surveillance zone.

The free status of the areas outside the containment zone would be suspended pending the establishment of the containment zone. The suspension of free status of these areas could be lifted reinstated irrespective of the provisions of Article 8.5.8., once the containment zone is clearly established, by complying with points 1 to 5 above. The containment zone should be managed in such a way that it can be demonstrated that commodities for international trade can be shown to have originated outside the containment zone.

The recovery of the FMD free status of the containment zone should follow the provisions of Article 8.5.8.

Recovery of free status

1. When a FMD outbreak or FMDV infection occurs in a FMD free country or zone where vaccination is not practised, one of the following waiting periods is required to regain the status of FMD free country or zone where vaccination is not practised:

   a) 3 months after the last case where a stamping-out policy and serological surveillance are applied in accordance with Articles 8.5.40. to 8.5.46.; or

   b) 3 months after the slaughter of all vaccinated animals where a stamping-out policy, emergency vaccination and serological surveillance are applied in accordance with Articles 8.5.40. to 8.5.46.; or

   c) 6 months after the last case or the last vaccination (according to the event that occurs the latest), where a stamping-out policy, emergency vaccination not followed by the slaughtering of all vaccinated animals, and serological surveillance are applied in accordance with Articles 8.5.40. to 8.5.46., provided that a serological survey based on the detection of antibodies to nonstructural proteins of FMDV demonstrates the absence of infection in the remaining vaccinated population.

   Where a stamping-out policy is not practised, the above waiting periods do not apply, and Article 8.5.2. or 8.5.4. applies.

2. When a FMD outbreak or FMDV infection occurs in a FMD free country or zone where vaccination is practised, one of the following waiting periods is required to regain the status of FMD free country or zone where vaccination is practised:

   a) 6 months after the last case where a stamping-out policy, emergency vaccination and serological surveillance in accordance with Articles 8.5.40. to 8.5.46. are applied, provided that the serological surveillance based on the detection of antibodies to nonstructural proteins of FMDV demonstrates the absence of virus circulation; or

   b) 18 months after the last case where a stamping-out policy is not applied, but emergency vaccination and serological surveillance in accordance with Articles 8.5.40. to 8.5.46. are applied, provided that the serological surveillance based on the detection of antibodies to nonstructural proteins of FMDV demonstrates the absence of virus circulation.
Article 8.5.9.

Transfer directly to slaughter of FMD susceptible animals from an infected zone to a free zone (where vaccination either is or is not practised) within a country

FMD susceptible animals should only leave the infected zone if moved by mechanised transport to the nearest designated abattoir located in the buffer protection zone directly to slaughter.

In the absence of an abattoir in the buffer protection zone, live FMD susceptible animals can be transported to the nearest abattoir in a free zone directly to slaughter only under the following conditions:

1. no FMD susceptible animal has been introduced into the establishment of origin and no animal in the establishment of origin has shown clinical signs of FMD for at least 30 days prior to movement;
2. the animals were kept in the establishment of origin for at least 3 months prior to movement;
3. FMD has not occurred within a 10-kilometre radius of the establishment of origin for at least 3 months prior to movement;
4. the animals must be transported under the supervision of the Veterinary Authority in a vehicle, which was cleansed and disinfected before loading, directly from the establishment of origin to the abattoir without coming into contact with other susceptible animals;
5. such an abattoir is not approved for the export of fresh meat during the time it is handling the meat of animals from the infected zone;
6. vehicles and the abattoir must be subjected to thorough cleansing and disinfection immediately after use.

All products obtained from the animals and any products coming into contact with them must be considered infected, and treated in such a way as to destroy any residual virus in accordance with Articles 8.5.32. to 8.5.39.

Animals moved into a free zone for other purposes must be moved under the supervision of the Veterinary Authority and comply with the conditions in Article 8.5.12.

Article 8.5.10.

Recommendations for importation from FMD free countries or zones where vaccination is not practised or FMD free zones where vaccination is not practised for FMD susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of FMD on the day of shipment;
2. were kept since birth or for at least the past 3 months in a FMD free country or zone where vaccination is not practised since birth or for at least the past 3 months;
3. have not been vaccinated.
Annex XVI (contd)

Article 8.5.11.

Recommendations for importation from FMD free countries or zones where vaccination is practised or from FMD-free zones where vaccination is practised for domestic ruminants and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of FMD on the day of shipment;
2. were kept in a FMD free country or zone since birth or for at least the past 3 months; and
3. have not been vaccinated and were subjected, with negative results, to tests for antibodies against FMD virus, when destined to a FMD free country or zone where vaccination is not practised.

Article 8.5.12.

Recommendations for importation from FMD infected countries or zones

for domestic ruminants and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of FMD on the day of shipment;
2. were kept in the establishment of origin since birth, or
   a) for the past 30 days, if a stamping-out policy is in force in the exporting country, or
   b) for the past 3 months, if a stamping-out policy is not in force in the exporting country, and that FMD has not occurred within a ten-kilometre radius of the establishment of origin for the relevant period as defined in points a) and b) above; and
3. were isolated in an establishment for the 30 days prior to shipment, and all animals in isolation were subjected to diagnostic tests (probang and serology) for evidence of FMDV infection with negative results at the end of that period, and that FMD did not occur within a ten-kilometre radius of the establishment during that period; or
4. were kept in a quarantine station for the 30 days prior to shipment, all animals in quarantine were subjected to diagnostic tests (probang and serology) for evidence of FMDV infection with negative results at the end of that period, and that FMD did not occur within a ten-kilometre radius of the quarantine station during that period;
5. were not exposed to any source of FMD infection during their transportation from the quarantine station to the place of shipment.

Article 8.5.13.

Recommendations for importation from FMD free countries or zones where vaccination is not practised or FMD-free zones where vaccination is not practised
for fresh semen of domestic ruminants and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of FMD on the day of collection of the semen;
   b) were kept for at least 3 months prior to collection in a FMD free country or zone where vaccination is not practised for at least 3 months prior to collection;

2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5. or Chapter 4.6., as relevant.

Article 8.5.14.

Recommendations for importation from FMD free countries or zones where vaccination is not practised or FMD free zones where vaccination is not practised for frozen semen of domestic ruminants and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of FMD on the day of collection of the semen and for the following 30 days;
   b) were kept in a FMD free country or zone where vaccination is not practised for at least 3 months prior to collection;

2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5. or Chapter 4.6., as relevant.

Article 8.5.15.

Recommendations for importation from FMD free countries or zones where vaccination is practised or from FMD free zones where vaccination is practised for semen of domestic ruminants and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of FMD on the day of collection of the semen and for the following 30 days;
   b) were kept for at least 3 months prior to collection in a country or zone free from FMD for at least 3 months prior to collection;
Annex XVI (contd)

c) if destined to a FMD free country or zone where vaccination is not practised:
   i) have not been vaccinated and were subjected, not less than 21 days after collection of the semen, to tests for antibodies against FMD virus, with negative results; or
   ii) had been vaccinated at least twice, with the last vaccination not more than 12 and not less than one month prior to collection;

2. no other animal present in the artificial insemination centre has been vaccinated within the month prior to collection;

3. the semen:
   a) was collected, processed and stored in conformity with the provisions of Chapter 4.5. or Chapter 4.6., as relevant;
   b) was stored in the country of origin for a period of at least one month following collection, and during this period no animal on the establishment where the donor animals were kept showed any sign of FMD.

Article 8.5.16.

Recommendations for importation from FMD infected countries or zones

for semen of domestic ruminants and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of FMD on the day of collection of the semen;
   b) were kept in an establishment where no animal had been added in the 30 days before collection, and that FMD has not occurred within 10 kilometres for the 30 days before and after collection;
   c) have not been vaccinated and were subjected, not less than 21 days after collection of the semen, to tests for antibodies against FMD virus, with negative results; or
   d) had been vaccinated at least twice, with the last vaccination not more than 12 and not less than one month prior to collection;

2. no other animal present in the artificial insemination centre has been vaccinated within the month prior to collection;

3. the semen:
   a) was collected, processed and stored in conformity with the provisions of Chapter 4.5. or Chapter 4.6., as relevant;
   b) was subjected, with negative results, to a test for FMDV infection if the donor animal has been vaccinated within the 12 months prior to collection;
c) was stored in the country of origin for a period of at least one month following collection, and during this period no animal on the establishment where the donor animals were kept showed any sign of FMD.

Article 8.5.17.

Recommendations for the importation of in vivo derived embryos of cattle

Irrespective of the FMD status of the exporting country or zone, Veterinary Authorities should authorise without restriction on account of FMD the import or transit through their territory of in vivo derived embryos of cattle subject to the presentation of an international veterinary certificate attesting that the embryos were collected, processed and stored in conformity with the provisions of Chapter 4.7 or Chapter 4.9.

Article 8.5.18.

Recommendations for importation from FMD free countries or zones where vaccination is not practised or FMD free zones where vaccination is not practised

for in vitro produced embryos of cattle

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) showed no clinical sign of FMD at the time of collection of the oocytes;
   b) were kept at the time of collection in a FMD free country or zone free from FMD where vaccination is not practised at the time of collection;

2. fertilisation was achieved with semen meeting the conditions referred to in Articles 8.5.13, 8.5.14, 8.5.15 or 8.5.16, as relevant;

3. the oocytes were collected, and the embryos were processed and stored in conformity with the provisions of Chapter 4.8 or Chapter 4.9, as relevant.

Article 8.5.19.

Recommendations for importation from FMD free countries or zones where vaccination is practised or from FMD free zones where vaccination is practised

for in vitro produced embryos of cattle

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) showed no clinical sign of FMD at the time of collection of the oocytes;
   b) were kept for at least 3 months prior to collection in a FMD free country or zone free from FMD where vaccination is practised for at least 3 months prior to collection;
Annex XVI (contd)

c) if destined for a FMD free country or zone where vaccination is not practised:

i) have not been vaccinated and were subjected, with negative results, to tests for antibodies against FMD virus; or

ii) had been vaccinated at least twice, with the last vaccination not less than one month and not more than 12 months prior to collection;

2. no other animal present in the establishment has been vaccinated within the month prior to collection;

3. fertilization was achieved with semen meeting the conditions referred to in Articles 8.5.13., 8.5.14., 8.5.15. or 8.5.16., as relevant;

4. the oocytes were collected, and the embryos were processed and stored in conformity with the provisions of Chapter 4.8. or Chapter 4.9., as relevant.

Article 8.5.20.

Recommendations for importation from FMD free countries or zones where vaccination is not practised or FMD free zones where vaccination is not practised for fresh meat of FMD susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which:

1. have been kept in the FMD free country or zone where vaccination is not practised since birth, or which have been imported in accordance with Article 8.5.10., Article 8.5.11. or Article 8.5.12.;

2. have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for FMD with favourable results.

Article 8.5.21.

Recommendations for importation from FMD free countries or zones where vaccination is practised or from FMD free zones where vaccination is practised for fresh meat of cattle and buffaloes (Bubalus bubalis) (excluding feet, head and viscera)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which:

1. have been kept in the FMD free country or zone where vaccination is practised since birth, or which have been imported in accordance with Article 8.5.10., Article 8.5.11. or Article 8.5.12.;

2. have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for FMD with favourable results.

Article 8.5.22.

Recommendations for importation from FMD free countries or zones where vaccination is practised or from FMD free zones where vaccination is practised for fresh meat or meat products of pigs and ruminants other than cattle and buffaloes
Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which:

1. have been kept in the FMD free country or zone where vaccination is practised since birth, or which have been imported in accordance with Article 8.5.10., Article 8.5.11. or Article 8.5.12.;

2. have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for FMD with favourable results.

Article 8.5.23.

Recommendations for importation from FMD infected countries or zones, where an official control programme exists, involving compulsory systematic vaccination of cattle

for fresh meat of cattle and buffaloes (Bubalus bubalis) (excluding feet, head and viscera)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat:

1. comes from animals which:
   a) have remained in the exporting country for at least 3 months prior to slaughter;
   b) have remained, during this period, in a part of the country where cattle are regularly vaccinated against FMD and where official controls are in operation;
   c) have been vaccinated at least twice with the last vaccination not more than 12 months and not less than one month prior to slaughter;
   d) were kept for the past 30 days in an establishment, and that FMD has not occurred within a ten-kilometre radius of the establishment during that period;
   e) have been transported, in a vehicle which was cleansed and disinfected before the cattle were loaded, directly from the establishment of origin to the approved abattoir without coming into contact with other animals which do not fulfil the required conditions for export;
   f) have been slaughtered in an approved abattoir:
      i) which is officially designated for export;
      ii) in which no FMD has been detected during the period between the last disinfection carried out before slaughter and the shipment for export has been dispatched;
   g) have been subjected to ante-mortem and post-mortem inspections for FMD with favourable results within 24 hours before and after slaughter;

2. comes from deboned carcasses:
   a) from which the major lymphatic nodes have been removed;
   b) which, prior to deboning, have been submitted to maturation at a temperature above + 2°C for a minimum period of 24 hours following slaughter and in which the pH value was below 6.0 when tested in the middle of both the longissimus dorsi.
Annex XVI (contd)

Article 8.5.24.

**Recommendations for importation from FMD infected countries or zones**

for meat products of domestic ruminants and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the entire consignment of meat comes from animals which have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for FMD with favourable results;

2. the meat has been processed to ensure the destruction of the FMD virus in conformity with one of the procedures referred to in Article 8.5.32.;

3. the necessary precautions were taken after processing to avoid contact of the meat products with any potential source of FMD virus.

Article 8.5.25.

**Recommendations for importation from FMD free countries or zones (where vaccination either is or is not practised)**

for milk and milk products intended for human consumption and for products of animal origin (from FMD susceptible animals) intended for use in animal feeding or for agricultural or industrial use

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products come from animals which have been kept in a FMD free country or zone since birth, or which have been imported in accordance with Article 8.5.10., Article 8.5.11. or Article 8.5.12.

Article 8.5.26.

**Recommendations for importation from FMD infected countries or zones where an official control programme exists**

for milk, cream, milk powder and milk products

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these products:
   a) originate from herds or flocks which were not infected or suspected of being infected with FMD at the time of milk collection;
   b) have been processed to ensure the destruction of the FMD virus in conformity with one of the procedures referred to in Article 8.5.36. and in Article 8.5.37.;

2. the necessary precautions were taken after processing to avoid contact of the products with any potential source of FMD virus.

Article 8.5.27.

**Recommendations for importation from FMD infected countries**
Annex XVI (contd)

for blood and meat-meals (from domestic or wild ruminants and pigs)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the manufacturing method for these products included heating to a minimum core temperature of 70°C for at least 30 minutes.

Article 8.5.28.

Recommendations for importation from FMD infected countries

for wool, hair, bristles, raw hides and skins (from domestic or wild ruminants and pigs)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these products have been processed to ensure the destruction of the FMD virus in conformity with one of the procedures referred to in Articles 8.5.33., 8.5.34. and 8.5.35.;

2. the necessary precautions were taken after collection or processing to avoid contact of the products with any potential source of FMD virus.

Veterinary Authorities can authorise, without restriction, the import or transit through their territory of semi-processed hides and skins (limed hides, pickled pelts, and semi-processed leather - e.g. wet blue and crust leather), provided that these products have been submitted to the usual chemical and mechanical processes in use in the tanning industry.

Article 8.5.29.

Recommendations for importation from FMD infected countries or zones

for straw and forage

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these commodities:

1. are free of grossly identifiable contamination with material of animal origin;

2. have been subjected to one of the following treatments, which, in the case of material sent in bales, has been shown to penetrate to the centre of the bale:

   a) either to the action of steam in a closed chamber such that the centre of the bales has reached a minimum temperature of 80°C for at least 10 minutes,

   b) or to the action of formalin fumes (formaldehyde gas) produced by its commercial solution at 35-40% in a chamber kept closed for at least 8 hours and at a minimum temperature of 19°C;

OR

3. have been kept in bond for at least 3 months (under study) before being released for export.

Article 8.5.30.

Recommendations for importation from FMD free countries or zones (where vaccination either is or is not practised)
Annex XVI (contd)

for skins and trophies derived from FMD susceptible wild animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products are derived from animals that have been killed in such a country or zone, or which have been imported from a country or zone free of FMD (where vaccination either is or is not practised).

Article 8.5.31.

Recommendations for importation from FMD infected countries or zones for skins and trophies derived from FMD susceptible wild animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products have been processed to ensure the destruction of the FMD virus in conformity with the procedures referred to in Article 8.5.38.

Article 8.5.32.

Procedures for the inactivation of the FMD virus in meat

For the inactivation of viruses present in meat, one of the following procedures should be used:

1. Canning

   Meat is subjected to heat treatment in a hermetically sealed container to reach an internal core temperature of at least 70°C for a minimum of 30 minutes or to any equivalent treatment which has been demonstrated to inactivate the FMD virus.

2. Thorough cooking

   Meat, previously deboned and defatted, shall be subjected to heating so that an internal temperature of 70°C or greater is maintained for a minimum of 30 minutes.

   After cooking, it shall be packed and handled in such a way that it cannot be exposed to a source of virus.

3. Drying after salting

   When rigor mortis is complete, the meat must be deboned, salted with cooking salt (NaCl) and completely dried. It must not deteriorate at ambient temperature.

   ‘Drying’ is defined in terms of the ratio between water and protein which must not be greater than 2.25:1.

Article 8.5.33.

Procedures for the inactivation of the FMD virus in wool and hair

For the inactivation of viruses present in wool and hair for industrial use, one of the following procedures should be used:

1. industrial washing, which consists of the immersion of the wool in a series of baths of water, soap and sodium hydroxide (soda) or potassium hydroxide (potash);
2. chemical depilation by means of slaked lime or sodium sulphide;

3. fumigation in formaldehyde in a hermetically sealed chamber for at least 24 hours. The most practical method is to place potassium permanganate in containers (which must NOT be made of plastic or polyethylene) and add commercial formalin; the amounts of formalin and potassium permanganate are respectively 53 ml and 35 g per cubic metre of the chamber;

4. industrial scouring which consists of the immersion of wool in a water-soluble detergent held at 60-70°C;

5. storage of wool at 18°C for 4 weeks, or 4°C for 4 months, or 37°C for 8 days.

Article 8.5.34.

Procedures for the inactivation of the FMD virus in bristles

For the inactivation of viruses present in bristles for industrial use, one of the following procedures should be used:

1. boiling for at least one hour;

2. immersion for at least 24 hours in a 1% solution of formaldehyde prepared from 30 ml commercial formalin per litre of water.

Article 8.5.35.

Procedures for the inactivation of the FMD virus in raw hides and skins

For the inactivation of viruses present in raw hides and skins for industrial use, the following procedure should be used: salting for at least 28 days in sea salt containing 2% sodium carbonate.

Article 8.5.36.

Procedures for the inactivation of the FMD virus in milk and cream for human consumption

For the inactivation of viruses present in milk and cream for human consumption, one of the following procedures should be used:

1. a sterilisation process applying a minimum temperature of 132°C for at least one second (ultra-high temperature [UHT]), or

2. if the milk has a pH less than 7.0, a sterilisation process applying a minimum temperature of 72°C for at least 15 seconds (high temperature - short time pasteurisation [HTST]), or

3. if the milk has a pH of 7.0 or over, the HTST process applied twice.

Article 8.5.37.

Procedures for the inactivation of the FMD virus in milk for animal consumption

For the inactivation of viruses present in milk for animal consumption, one of the following procedures should be used:

1. the HTST process applied twice;
Annex XVI (contd)

2. HTST combined with another physical treatment, e.g. maintaining a pH 6 for at least one hour or additional heating to at least 72°C combined with dessication;

3. UHT combined with another physical treatment referred to in point 2 above.

Article 8.5.38.

Procedures for the inactivation of the FMD virus in skins and trophies from wild animals susceptible to the disease

For the inactivation of viruses present in skins and trophies from wild animals susceptible to FMD, one of the following procedures should be used prior to complete taxidermal treatment:

1. boiling in water for an appropriate time so as to ensure that any matter other than bone, horns, hooves, claws, antlers or teeth is removed;

2. gamma irradiation at a dose of at least 20 kiloGray at room temperature (20°C or higher);

3. soaking, with agitation, in a 4% (w/v) solution of washing soda (sodium carbonate - Na₂CO₃) maintained at pH 11.5 or above for at least 48 hours;

4. soaking, with agitation, in a formic acid solution (100 kg salt [NaCl] and 12 kg formic acid per 1,000 litres water) maintained at below pH 3.0 for at least 48 hours; wetting and dressing agents may be added;

5. in the case of raw hides, salting for at least 28 days with sea salt containing 2% washing soda (sodium carbonate - Na₂CO₃).

Article 8.5.39.

Procedures for the inactivation of the FMD virus in casings of small ruminants and pigs

For the inactivation of viruses present in casings of small ruminants and pigs, the following procedures should be used: salting for at least 30 days either with dry salt (NaCl) or with saturated brine (Aw < 0.80), or with phosphate salts/sodium chloride mixture, and kept at room temperature at about 20°C during this entire period.

Article 8.5.40.

Surveillance: introduction

Articles 8.5.40. to 8.5.46. define the principles and provide a guide for the surveillance of FMD in accordance with Chapter 1.4. applicable to Members seeking recognition from the OIE for establishment of freedom from FMD, either with or without the use of vaccination. This may be for the entire country or a zone within the country. Guidance is provided for Members seeking reestablishment of freedom from FMD for the whole entire country or for a zone within the country, either with or without vaccination, following an outbreak, as well as recommendations and for the maintenance of FMD status are provided. Applications to the OIE for recognition of freedom should follow the format and answer all the questions posed by the “Questionnaire on FMD” available from the OIE Central Bureau.
The impact and epidemiology of FMD differ widely in different regions of the world and therefore it is impossible to provide specific recommendations for all situations. It is axiomatic that the surveillance strategies employed for demonstrating freedom from FMD at an acceptable level of confidence will need to be adapted to the local situation. For example, the approach to proving freedom from FMD following an outbreak caused by a pig-adapted strain of FMD virus (FMDV) should differ significantly from an application designed to prove freedom from FMD for a country or zone where African buffaloes (Syncerus caffer) provide a potential reservoir of infection. It is incumbent upon the Member to submit a dossier to the OIE in support of its application that not only explains the epidemiology of FMD in the region concerned but also demonstrates how all the risk factors are managed. This should include provision of scientifically-based supporting data. There is therefore considerable latitude available to Members to provide a well-reasoned argument to prove that the absence of FMDV infection (in non-vaccinated populations) or circulation (in vaccinated populations) is assured at an acceptable level of confidence.

Surveillance for FMD should be in the form of a continuing programme designed to establish that the whole territory or part of it is free from FMDV infection/circulation.

For the purposes of this Chapter, virus circulation means transmission of FMDV as demonstrated by clinical signs, serological evidence or virus isolation.

Article 8.5.41.

**Surveillance: general conditions and methods**

1. A surveillance system in accordance with Chapter 1.4. should be under the responsibility of the Veterinary Authority. A procedure should be in place for the rapid collection and transport of samples from suspect cases of FMD to a laboratory for FMD diagnoses as described in the Terrestrial Manual.

2. The FMD surveillance programme should:

   a) include an early warning system throughout the production, marketing and processing chain for reporting suspicious cases. Farmers and workers who have day-to-day contact with livestock, as well as diagnosticians, should report promptly any suspicion of FMD. They should be supported directly or indirectly (e.g. through private veterinarians or veterinary para-professionals) by government information programmes and the Veterinary Authority. All suspect cases of FMD should be investigated immediately. Where suspicion cannot be resolved by epidemiological and clinical investigation, samples should be taken and submitted to a laboratory. This requires that sampling kits and other equipment are available for those responsible for surveillance. Personnel responsible for surveillance should be able to call for assistance from a team with expertise in FMD diagnosis and control;

   b) implement, when relevant, regular and frequent clinical inspection and serological testing of high-risk groups of animals, such as those adjacent to a FMD infected country or infected zone (for example, bordering a game park in which infected wildlife are present).

An effective surveillance system will periodically identify suspicious cases that require follow-up and investigation to confirm or exclude that the cause of the condition is FMDV. The rate at which such suspicious cases are likely to occur will differ between epidemiological situations and cannot therefore be predicted reliably. Applications for freedom from FMDV infection/circulation should, in consequence, provide details of the occurrence of suspicious cases and how they were investigated and dealt with. This should include the results of laboratory testing and the control measures to which the animals concerned were subjected during the investigation (quarantine, movement stand-still orders, etc.).
Annex XVI (contd)

Article 8.5.42.

Surveillance strategies

1. Introduction

The target population for surveillance aimed at identifying disease and infection should cover all the susceptible species within the country or zone to be recognised as free from FMDV infection/circulation.

The design of surveillance programmes to prove the absence of FMDV infection/circulation needs to be carefully followed to avoid producing results that are either insufficiently reliable to be accepted by the OIE or international trading partners, or excessively costly and logistically complicated. The design of any surveillance programme, therefore, requires inputs from professionals competent and experienced in this field.

The strategy employed may be based on randomised sampling requiring surveillance consistent with demonstrating the absence of FMDV infection/circulation at an acceptable level of statistical confidence. The frequency of sampling should be dependent on the epidemiological situation.

Targeted surveillance (e.g. based on the increased likelihood of infection in particular localities or species) may be an appropriate strategy. The Member should justify the surveillance strategy chosen as adequate to detect the presence of FMDV infection/circulation in accordance with Chapter 1.4. and the epidemiological situation. It may, for example, be appropriate to target clinical surveillance at particular species likely to exhibit clear clinical signs (e.g. cattle and pigs). If a Member wishes to apply for recognition of a specific zone within the country as being free from FMDV infection/circulation, the design of the survey and the basis for the sampling process would need to be aimed at the population within the zone.

For random surveys, the design of the sampling strategy will need to incorporate an epidemiologically appropriate design prevalence. The sample size selected for testing will need to be large enough to detect infection/circulation if it were to occur at a predetermined minimum rate. The sample size and expected disease prevalence determine the level of confidence in the results of the survey. The Member must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Chapter 1.4. Selection of the design prevalence in particular clearly needs to be based on the prevailing or historical epidemiological situation.

Irrespective of the survey design selected, the sensitivity and specificity of the diagnostic tests employed are key factors in the design, sample size determination and interpretation of the results obtained. Ideally, the sensitivity and specificity of the tests used should be validated for the vaccination/infection history and production class of animals in the target population.

Irrespective of the testing system employed, surveillance design should anticipate the occurrence of false positive reactions. If the characteristics of the testing system are known, the rate at which these false positives are likely to occur can be calculated in advance. There needs to be an effective procedure for following-up positives to ultimately determine with a high level of confidence, whether they are indicative of infection/circulation or not. This should involve both supplementary tests and follow-up investigation to collect diagnostic material from the original sampling unit as well as herds which may be epidemiologically linked to it.

The principles involved in surveillance for disease/infection are technically well defined. The design of surveillance programmes to prove the absence of FMDV infection/circulation needs to be carefully followed to avoid producing results that are either insufficiently reliable to be accepted by the OIE or international trading partners, or excessively costly and logistically complicated. The design of any surveillance programme, therefore, requires inputs from professionals competent and experienced in this field.
2. **Clinical surveillance**

Clinical surveillance aims at detecting clinical signs of FMD by close physical examination of susceptible animals. Whereas significant emphasis is placed on the diagnostic value of mass serological screening, surveillance based on clinical inspection should not be underrated. It may be able to provide a high level of confidence of detection of disease if a sufficiently large number of clinically susceptible animals is examined.

Clinical surveillance and laboratory testing should always be applied in series to clarify the status of FMD suspects detected by either of these complementary diagnostic approaches. Laboratory testing may confirm clinical suspicion, while clinical surveillance may contribute to confirmation of positive serology. Any sampling unit within which suspicious animals are detected should be classified as infected until contrary evidence is produced.

A number of issues must be considered in clinical surveillance for FMD. The often underestimated labour intensity and the logistical difficulties involved in conducting clinical examinations should not be underestimated and should be taken into account.

Identification of clinical cases is fundamental to FMD surveillance. Establishment of the molecular, antigenic and other biological characteristics of the causative virus, as well as its source, is dependent upon disclosure of such animals. It is essential that FMDV isolates are sent regularly to the regional reference laboratory for genetic and antigenic characterization.

3. **Virological surveillance**

Virological surveillance using tests described in the Terrestrial Manual should be conducted:

a) to monitor at risk populations;

b) to confirm clinically suspect cases;

c) to follow up positive serological results;

d) to test “normal” daily mortality, to ensure early detection of infection in the face of vaccination or in establishments epidemiologically linked to an outbreak.

4. **Serological surveillance**

Serological surveillance aims at detecting antibodies against FMDV. Positive FMDV antibody test results can have four possible causes:

a) natural infection with FMDV;

b) vaccination against FMD;

c) maternal antibodies derived from an immune dam (maternal antibodies in cattle are usually found only up to 6 months of age but in some individuals and in some species, maternal antibodies can be detected for considerably longer periods);

d) heterophile (cross) reactions.
Annex XVI (contd)

It is important that serological tests, where applicable, contain antigens appropriate for detecting antibodies against viral variants (types, subtypes, lineages, topotypes, etc.) that have recently occurred in the region concerned. Where the probable identity of FMDVs is unknown or where exotic viruses are suspected to be present, tests able to detect representatives of all serotypes should be employed (e.g. tests based on nonstructural viral proteins – see below).

It may be possible to use serum collected for other survey purposes for FMD surveillance. However, the principles of survey design described in this Chapter and the requirement for a statistically valid survey for the presence of FMDV should not be compromised.

The discovery of clustering of seropositive reactions should be foreseen. It may reflect any of a series of events, including but not limited to the demographics of the population sampled, vaccinal exposure or the presence of field strain infection. As clustering may signal field strain infection, the investigation of all instances must be incorporated in the survey design. If vaccination cannot be excluded as the cause of positive serological reactions, diagnostic methods should be employed that detect the presence of antibodies to nonstructural proteins (NSPs) of FMDVs as described in the Terrestrial Manual.

The results of random or targeted serological surveys are important in providing reliable evidence that FMDV infection is not present in a country or zone. It is therefore essential that the survey be thoroughly documented.

Article 8.5.43.

Members applying for recognition of freedom from FMD for the whole country or a zone where vaccination is not practised: additional surveillance procedures

In addition to the general conditions described in the above-mentioned articles, a Member applying for recognition of FMD freedom for the country or a zone where vaccination is not practised should provide evidence for the existence of an effective surveillance programme. The strategy and design of the surveillance programme will depend on the prevailing epidemiological circumstances and will be planned and implemented according to general conditions and methods in this Chapter, to demonstrate absence of FMDV infection, during the preceding 12 months in susceptible populations. This requires the support of a national or other laboratory able to undertake identification of FMDV infection through virus/antigen/genome detection and antibody tests described in the Terrestrial Manual.

Article 8.5.44.

Members applying for recognition of freedom from FMD for the whole country or a zone where vaccination is practised: additional surveillance procedures

In addition to the general conditions described in the above-mentioned articles, a Member applying for recognition of country or zone freedom from FMD with vaccination should show evidence of an effective surveillance programme planned and implemented according to general conditions and methods in this Chapter. Absence of clinical disease in the country or zone for the past 2 years should be demonstrated. Furthermore, surveillance should demonstrate that FMDV has not been circulating in any susceptible population during the past 12 months. This will require serological surveillance incorporating tests able to detect antibodies to NSPs as described in the Terrestrial Manual. Vaccination to prevent the transmission of FMDV may be part of a disease control programme. The level of herd immunity required to prevent transmission will depend on the size, composition (e.g. species) and density of the susceptible population. It is therefore impossible to be prescriptive. However, the aim should, in general, be to vaccinate at least 80% of the susceptible population. The vaccine must comply with the Terrestrial Manual. Based on the epidemiology of FMD in the country or zone, it may be that a decision is reached to vaccinate only certain species or other subsets of the total susceptible population. In that case, the rationale should be contained within the dossier accompanying the application to the OIE for recognition of status.
Evidence to show the effectiveness of the vaccination programme should be provided.

Article 8.5.45.

Members re-applying for recognition of freedom from FMD for the whole country or a zone where vaccination is either practised or not practised, following an outbreak: additional surveillance procedures

In addition to the general conditions described in the above-mentioned articles, a country re-applying for country or zone freedom from FMD where vaccination is practised or not practised should show evidence of an active surveillance programme for FMD as well as absence of FMDV infection/circulation.

This will require serological surveillance incorporating, in the case of a country or a zone practising vaccination, tests able to detect antibodies to NSPs as described in the Terrestrial Manual.

Four strategies are recognised by the OIE in a programme to eradicate FMDV infection following an outbreak:

1. slaughter of all clinically affected and in-contact susceptible animals;
2. slaughter of all clinically affected and in-contact susceptible animals and vaccination of at-risk animals, with subsequent slaughter of vaccinated animals;
3. slaughter of all clinically affected and in-contact susceptible animals and vaccination of at-risk animals, without subsequent slaughter of vaccinated animals;
4. vaccination used without slaughter of affected animals or subsequent slaughter of vaccinated animals.

The time periods before which an application can be made for reinstatement of freedom from FMD depends on which of these alternatives is followed. The time periods are prescribed in Article 8.5.8.

In all circumstances, a Member re-applying for country or zone freedom from FMD with vaccination or without vaccination should report the results of an active surveillance programme implemented according to general conditions and methods in this Chapter.

Article 8.5.46.

The use and interpretation of serological tests (see Figure 1)

The recommended serological tests for FMD surveillance are described in the Terrestrial Manual.

Animals infected with FMDV produce antibodies to both the structural proteins (SP) and the nonstructural proteins (NSP) of the virus. Tests for SP antibodies to include SP-ELISAs and the virus neutralisation test (VNT). The SP tests are serotype specific and for optimal sensitivity should utilise an antigen or virus closely related to the field strain against which antibodies are being sought. Tests for NSP antibodies include NSP I ELISA 3ABC and the electro-immunotransfer blotting technique (EITB) as recommended in the Terrestrial Manual or equivalent validated tests. In contrast to SP tests, NSP tests can detect antibodies to all serotypes of FMD virus. Animals vaccinated and subsequently infected with FMD virus develop antibodies to NSPs, but in some, the titre may be lower than that found in infected animals that have not been vaccinated. Both the NSP I ELISA 3ABC and EITB tests have been extensively used in cattle. Validation in other species is ongoing. Vaccines used should comply with the standards of the Terrestrial Manual insofar as purity is concerned to avoid interference with NSP antibody testing.
Annex XVI (contd)

Serological testing is a suitable tool for FMD surveillance. The choice of a serosurveillance system will depend on, amongst other things, the vaccination status of the country. A country, which is free from FMD without vaccination, may choose serosurveillance of high-risk subpopulations (e.g. based on geographical risk for exposure to FMDV). SP tests may be used in such situations for screening sera for evidence of FMDV infection/circulation if a particular virus of serious threat has been identified and is well characterised. In other cases, NSP testing is recommended in order to cover a broader range of strains and even serotypes. In both cases, serological testing can provide additional support to clinical surveillance. Regardless of whether SP or NSP tests are used in countries that do not vaccinate, a diagnostic follow-up protocol should be in place to resolve any presumptive positive serological test results.

In areas where animals have been vaccinated, SP antibody tests may be used to monitor the serological response to the vaccination. However, NSP antibody tests should be used to monitor for FMDV infection/circulation. NSP-ELISAs may be used for screening sera for evidence of infection/circulation irrespective of the vaccination status of the animal. All herds with seropositive reactors should be investigated. Epidemiological and supplementary laboratory investigation results should document the status of FMDV infection/circulation for each positive herd. Tests used for confirmation should be of high diagnostic specificity to eliminate as many false positive screening test reactors as possible. The diagnostic sensitivity of the confirmatory test should approach that of the screening test. The EITB or another OIE-accepted test should be used for confirmation.

Information should be provided on the protocols, reagents, performance characteristics and validation of all tests used.

1. The follow-up procedure in case of positive test results if no vaccination is used in order to establish or re-establish FMD free status without vaccination

Any positive test result (regardless of whether SP or NSP tests were used) should be followed up immediately using appropriate clinical, epidemiological, serological and, where possible, virological investigations of the reactor animal at hand, of susceptible animals of the same epidemiological unit and of susceptible animals that have been in contact or otherwise epidemiologically associated with the reactor animal. If the follow-up investigations provide no evidence for FMDV infection, the reactor animal shall be classified as FMD negative. In all other cases, including the absence of such follow-up investigations, the reactor animal should be classified as FMD positive.

2. The follow-up procedure in case of positive test results if vaccination is used in order to establish or re-establish FMD free status with vaccination

In case of vaccinated populations, one has to exclude that positive test results are indicative of virus circulation. To this end, the following procedure should be followed in the investigation of positive serological test results derived from surveillance conducted on FMD vaccinated populations.

The investigation should examine all evidence that might confirm or refute the hypothesis that the positive results to the serological tests employed in the initial survey were not due to virus circulation.

All the epidemiological information should be substantiated, and the results should be collated in the final report.

It is suggested that in the primary sampling units where at least one animal reacts positive to the NSP test, the following strategy(ies) should be applied:
a) Following clinical examination, a second serum sample should be taken from the animals tested in the initial survey after an adequate interval of time has lapsed, on the condition that they are individually identified, accessible and have not been vaccinated during this period. Antibody titres against NSP at the time of retest should be statistically either equal to or lower than those observed in the initial test if virus is not circulating.

The animals sampled should remain in the holding pending test results and should be clearly identifiable. If the three conditions for retesting mentioned above cannot be met, a new serological survey should be carried out in the holding after an adequate period of time, repeating the application of the primary survey design and ensuring that all animals tested are individually identified. These animals should remain in the holding and should not be vaccinated, so that they can be retested after an adequate period of time.

b) Following clinical examination, serum samples should be collected from representative numbers of cattle that were in physical contact with the primary sampling unit. The magnitude and prevalence of antibody reactivity observed should not differ in a statistically significant manner from that of the primary sample if virus is not circulating.

c) Following clinical examination, epidemiologically linked herds should be serologically tested and satisfactory results should be achieved if virus is not circulating.

d) Sentinel animals can also be used. These can be young, unvaccinated animals or animals in which maternally conferred immunity has lapsed and belonging to the same species resident within the positive initial sampling units. They should be serologically negative if virus is not circulating. If other susceptible, unvaccinated ruminants (sheep, goats) are present, they could act as sentinels to provide additional serological evidence.

Laboratory results should be examined in the context of the epidemiological situation. Corollary information needed to complement the serological survey and assess the possibility of viral circulation includes but is not limited to:

- characterization of the existing production systems;

- results of clinical surveillance of the suspects and their cohorts;

- quantification of vaccinations performed on the affected sites;

- sanitary protocol and history of the establishments with positive reactors;

- control of animal identification and movements;

- other parameters of regional significance in historic FMDV transmission.

The entire investigative process should be documented as standard operating procedure within the surveillance programme.
Annex XVI (contd)

Fig. 1. Schematic representation of laboratory tests for determining evidence of FMDV infection through or following serological surveys

Key:

<table>
<thead>
<tr>
<th>Test</th>
<th>Description</th>
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<tbody>
<tr>
<td>ELISA</td>
<td>Enzyme-linked immunosorbent assay</td>
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<tr>
<td>VNT</td>
<td>Virus neutralisation test</td>
</tr>
<tr>
<td>NSP</td>
<td>Nonstructural protein(s) of foot and mouth disease virus (FMDV)</td>
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<tr>
<td>3ABC</td>
<td>NSP antibody test</td>
</tr>
<tr>
<td>EITB</td>
<td>Electro-immuno transfer blotting technique (Western blot for NSP antibodies of FMDV)</td>
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<tr>
<td>SP</td>
<td>Structural protein test</td>
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<tr>
<td>S</td>
<td>No evidence of FMDV</td>
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</table>
CHAPTER 8.8.

LEPTOSPIROSIS

Article 8.8.1.

Under study.

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CHAPTER 8.11.

RABIES

Article 8.11.1.

General provisions

For the purposes of the Terrestrial Code, the incubation period for rabies shall be 6 months, and the infective period in domestic carnivores starts 15 days before the onset of the first clinical signs and ends when the animal dies.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 8.11.2.

Rabies free country

For the purposes of international trade, a country may be considered free from rabies when:

1. the disease is notifiable;
2. an effective system of disease surveillance is in operation;
3. all regulatory measures for the prevention and control of rabies have been implemented including effective importation procedures;
4. no case of indigenously acquired rabies infection has been confirmed in man or any animal species during the past 2 years; however, this status would not be affected by the isolation of an Australian or European Bats Lyssavirus;
5. no imported case in carnivores has been confirmed outside a quarantine station for the past 6 months.

Article 8.11.3.

Recommendations for importation from rabies free countries

for domestic mammals, and wild mammals reared under confined conditions

Veterinary authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of rabies on the day of shipment;
2. were kept since birth or for the 6 months prior to shipment in a rabies free country or were imported in conformity with the regulations stipulated in Articles 8.11.5., 8.11.6. or 8.11.7.

Article 8.11.4.

Recommendations for importation from rabies free countries

for wild mammals not reared under confined conditions
Annex XVIII (contd)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of rabies on the day of shipment;

2. have been captured in a rabies free country, at a sufficient distance from any infected country. The distance should be defined according to the species exported and the reservoir species in the infected country.

Article 8.11.5.

Recommendations for importation from countries considered infected with rabies

for dogs and cats

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of rabies within 48 hours of shipment;

AND EITHER

2. were identified by a permanent mark (such as a microchip) and their identification number shall be stated in the certificate; and

3. were vaccinated against rabies:
   a) not less than 6 months and not more than one year prior to shipment in the case of a primary vaccination, which should have been carried out when the animals were at least 3 months old;
   b) not more than one year prior to shipment in the case of a booster vaccination;
   c) with an inactivated virus vaccine or with a recombinant vaccine expressing the rabies virus glycoprotein; and

4. were subjected not less than 3 months and not more than 24 months prior to shipment to an antibody test as prescribed in the Terrestrial Manual with a positive result equivalent to at least 0.5 IU/ml;

OR

5. have not been vaccinated against rabies or do not meet all the conditions set out in points 2, 3 and 4 above; in such cases, the importing country may require the placing of the animals in a quarantine station located on its territory, in conformity with the conditions stipulated in its animal health legislation.

Article 8.11.6.

Recommendations for importation from countries considered infected with rabies

for domestic ruminants, equines and pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of rabies on the day of shipment;
2. were kept for the 6 months prior to shipment in an establishment where separation from wild and feral animals was maintained and where no case of rabies was reported for at least 12 months prior to shipment.

Article 8.11.7.

Recommendations for importation from countries considered infected with rabies

for laboratory reared rodents and lagomorphs, and lagomorphs or wild mammals (other than non-human primates) reared under confined conditions

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of rabies on the day of shipment;
2. were kept since birth, or for the 12 months prior to shipment, in an establishment where no case of rabies was reported for at least 12 months prior to shipment.

Article 8.11.8.

Recommendations for importation from countries considered infected with rabies

for wild mammals not belonging to the orders of primates or carnivores and not reared under confined conditions

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of rabies on the day of shipment;
2. were kept in a quarantine station for the 6 months prior to shipment.

Article 8.11.9.

Recommendations for importation from countries considered infected with rabies

for frozen semen of dogs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the donor animals showed no clinical sign of rabies during the 15 days following collection of the semen.

[Note: For non-human primates, reference should be made to Chapter 6.9.]
CHAPTER 8.13.

RINDERPEST

Article 8.13.1.

General provisions

For the purposes of the Terrestrial Code, the incubation period for rinderpest (RP) shall be 21 days.

For the purpose of this Chapter, a case includes an animal infected with rinderpest virus (RPV).

For the purpose of this Chapter, susceptible animals apply to both domestic and wild artiodactyls.

For the purposes of international trade, this Chapter deals not only with the occurrence of clinical signs caused by RPV, but also with the presence of infection with RPV in the absence of clinical signs.

Ban on vaccination against rinderpest means a ban on administering a RP vaccine to any susceptible animal and a heterologous vaccine against RP to any large ruminants or pigs.

1. Animal not vaccinated against RP means:
   a) for large ruminants and pigs: an animal that has received neither a RP vaccine nor a heterologous vaccine against RP;
   b) for small ruminants: an animal that has not received a RP vaccine.

2. The following defines the occurrence of RPV infection:
   a) RPV has been isolated and identified as such from an animal or a product derived from that animal; or
   b) viral antigen or viral ribonucleic acid (RNA) specific to RP has been identified in samples from one or more animals showing one or more clinical signs consistent with RP, or epidemiologically linked to an outbreak of RP, or giving cause for suspicion of association or contact with RP; or
   c) antibodies to RPV antigens which are not the consequence of vaccination, have been identified in one or more animals with either epidemiological links to a confirmed or suspected outbreak of RP in susceptible animals, or showing clinical signs consistent with recent infection with RP.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 8.13.2.

Rinderpest free country

To qualify for inclusion in the existing list of RP free countries, a Member should:

1. have a record of regular and prompt animal disease reporting:
Annex XIX (contd)

2. send a declaration to the OIE stating that:
   a) there has been no outbreak of RP during the past 24 months,
   b) no evidence of RPV infection has been found during the past 24 months,
   c) no vaccination against RP has been carried out during the past 24 months,

3. and supply documented evidence that surveillance for both RP and RPV infection in accordance with Articles 8.13.20. to 8.13.27. is in operation and that regulatory measures for the prevention and control of RP have been implemented;

4. not have imported since the cessation of vaccination any animals vaccinated against RP.

The Member will be included in the list only after the submitted evidence has been accepted by the OIE. Retention on the list requires that the information in points 2a), 2b), 2c), and 3 above be re-submitted annually and changes in the epidemiological situation or other significant events should be reported to the OIE according to the requirements in Chapter 1.1.

Article 8.13.3.

Recovery of free status

When a RP outbreak or RPV infection occurs in a RP free country, one of the following waiting periods is required to regain the status of RP free country:

1. 3 months after the last case where a stamping-out policy and serological surveillance are applied in accordance with Articles 8.13.20. to 8.13.27.; or

2. 3 months after the slaughter of all vaccinated animals where a stamping-out policy, emergency vaccination and serological surveillance are applied in accordance with Articles 8.13.20. to 8.13.27.; or

3. 6 months after the last case or the last vaccination (according to the event that occurs the latest), where a stamping-out policy, emergency vaccination not followed by the slaughter of all vaccinated animals, and serological surveillance are applied in accordance with Articles 8.13.20. to 8.13.27.

Where a stamping-out policy is not practised, the above waiting periods do not apply but Article 8.13.2. applies.

Article 8.13.4.

Infected country

When the requirements for acceptance as a RP free country are not fulfilled, a country shall be considered as RP infected.

Article 8.13.5.

Recommendations for importation from RP free countries

for RP susceptible animals
Veterinary authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of RP on the day of shipment;
2. remained in a RP free country since birth or for at least 30 days prior to shipment.

**Article 8.13.6.**

**Recommendations for importation from RP infected countries**

for RP susceptible animals

Veterinary authorities should require the presentation of an international veterinary certificate attesting that:

1. RP is the subject of a national surveillance programme according to Articles 8.13.20. to 8.13.27.;
2. RP has not occurred within a 10-kilometre radius of the establishment of origin of the animals destined for export for at least 21 days prior to their shipment to the quarantine station referred to in point 3b) below;
3. the animals:
   a) showed no clinical sign of RP on the day of shipment;
   b) were kept in the establishment of origin since birth or for at least 21 days before introduction into the quarantine station referred to in point c) below;
   c) have not been vaccinated against RP, were isolated in a quarantine station for the 30 days prior to shipment, and were subjected to a diagnostic test for RP on two occasions with negative results, at an interval of not less than 21 days;
   d) were not exposed to any source of infection during their transportation from the quarantine station to the place of shipment;
4. RP has not occurred within a ten-kilometre radius of the quarantine station for 30 days prior to shipment.

**Article 8.13.7.**

**Recommendations for importation from RP free countries**

for semen of RP susceptible animals

Veterinary authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of RP on the day of collection of the semen;
   b) were kept in a RP free country for at least 3 months prior to collection;
2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5.
Annex XIX (contd)

Article 8.13.8.

**Recommendations for importation from RP infected countries**

for semen of RP susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. RP is the subject of a national surveillance programme according to Articles 8.13.20. to 8.13.27.;

2. the donor animals:
   
   a) showed no clinical sign of RP on the day of collection of the semen;
   
   b) were kept in an establishment where no RP susceptible animals had been added in the 21 days before collection, and that RP has not occurred within 10 kilometres of the establishment for the 21 days before and after collection;
   
   c) were vaccinated against RP at least 3 months prior to collection; or
   
   d) have not been vaccinated against RP, and were subjected to a diagnostic test on two occasions with negative results, at an interval of not less than 21 days within the 30 days prior to collection;

3. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5.

Article 8.13.9.

**Recommendations for importation from RP free countries**

for in vivo derived embryos of RP susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor females were kept in an establishment located in a RP free country at the time of collection;

2. the embryos were collected, processed and stored in conformity with the provisions of Chapter 4.7.

Article 8.13.10.

**Recommendations for importation from RP infected countries**

for in vivo derived embryos of RP susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. RP is the subject of a national surveillance programme according to Articles 8.13.20. to 8.13.27.;

2. the donor females:
   
   a) and all other animals in the establishment showed no clinical sign of RP at the time of collection and for the following 21 days;
b) were kept in an establishment where no RP susceptible animals had been added in the 21 days before collection of the embryos;

c) were vaccinated against RP at least 3 months prior to collection; or

d) have not been vaccinated against RP, and were subjected to a diagnostic test for RP on two occasions with negative results, at an interval of not less than 21 days within the 30 days prior to collection;

3. the embryos were collected, processed and stored in conformity with the provisions of Chapter 4.7.

Article 8.13.11.

Recommendations for importation from RP free countries

for fresh meat or meat products of susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment comes from animals which have been kept in the country since birth or for at least 3 months prior to slaughter.

Article 8.13.12.

Recommendations for importation from RP infected countries

for fresh meat (excluding offal) of susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat:

1. comes from a country where RP is the subject of a national surveillance programme according to Articles 8.13.20. to 8.13.27.;

2. comes from animals which:

   a) showed no clinical sign of RP within 24 hours before slaughter;

   b) have remained in the country for at least 3 months prior to slaughter;

   c) were kept in the establishment of origin since birth or for at least 30 days prior to shipment to the approved abattoir, and that RP has not occurred within a ten-kilometre radius of the establishment during that period;

   d) were vaccinated against RP at least 3 months prior to shipment to the approved abattoir;

   e) had been transported, in a vehicle which was cleansed and disinfected before the animals were loaded, directly from the establishment of origin to the approved abattoir without coming into contact with other animals which do not fulfil the required conditions for export;

   f) were slaughtered in an approved abattoir in which no RP has been detected during the period between the last disinfection carried out before abattoir and the date on which the shipment has been dispatched.
Article 8.13.13.

**Recommendations for importation from RP infected countries**

for meat products of susceptible animals

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. only fresh meat complying with the provisions of Article 8.13.12. has been used in the preparation of the meat products or

2. the meat products have been processed to ensure the destruction of the RPV in conformity with one of the procedures referred to in Article 8.5.32.;

3. the necessary precautions were taken after processing to avoid contact of the meat products with any possible source of RPV.


**Recommendations for importation from RP free countries**

for milk and milk products intended for human consumption and for products of animal origin (from RP susceptible animals) intended for use in animal feeding or for agricultural or industrial use

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products come from animals which have been kept in the country since birth or for at least 3 months.

Article 8.13.15.

**Recommendations for importation from RP infected countries**

for milk and cream

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these products:
   a) originate from herds or flocks which were not subjected to any restrictions due to RP at the time of milk collection;

   b) have been processed to ensure the destruction of the RPV in conformity with one of the procedures referred to in Articles 8.5.36. and 8.5.37.;

2. the necessary precautions were taken after processing to avoid contact of the products with any potential source of RPV.

Article 8.13.16.

**Recommendations for importation from RP infected countries**

for milk products
Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these products are derived from milk complying with the above requirements;
2. the necessary precautions were taken after processing to avoid contact of the milk products with a potential source of RPV.

Article 8.13.17.

**Recommendations for importation from RP infected countries**

for blood and meat-meals (from susceptible animals)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the manufacturing method for these products included heating to a minimum internal temperature of 70°C for at least 30 minutes.

Article 8.13.18.

**Recommendations for importation from RP infected countries**

for wool, hair, bristles, raw hides and skins (from susceptible animals)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these products have been processed to ensure the destruction of the RPV in conformity with one of the procedures referred to in Articles 8.5.33., 8.5.34. and 8.5.35.;
2. the necessary precautions were taken after processing to avoid contact of the products with any potential source of RPV.

Veterinary Authorities can authorise, without restriction, the import or transit through their territory of semi-processed hides and skins (limed hides, pickled pelts, and semi-processed leather - e.g. wet blue and crust leather), provided that these products have been submitted to the usual chemical and mechanical processes in use in the tanning industry.

Article 8.13.19.

**Recommendations for importation from RP infected countries**

for hooves, claws, bones and horns, hunting trophies and preparations destined for museums (from susceptible animals)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products:

1. were completely dried and had no trace on them of skin, flesh or tendon; and/or
2. have been adequately disinfected.
Annex XIX (contd)

Article 8.13.20.

Surveillance: introduction

In order to receive OIE recognition of rinderpest freedom, a country’s national authority must present for consideration a dossier of information relating to its livestock production systems, rinderpest vaccination and eradication history and the functioning of its Veterinary Services. The dossier must contain convincing evidence derived from an animal disease surveillance system that sufficient evidence has accrued to demonstrate that the presence of rinderpest virus would have been disclosed were it to be present. Recommendations on the structure and the functioning of Veterinary Services and diagnostic support services are provided in Chapters 3.1. and 3.2. of the Terrestrial Code. A Member must also be in compliance with its OIE reporting obligations (Chapter 1.1. of the Terrestrial Code).

Articles 8.13.20. to 8.13.27. define the principles and provides a guide for the surveillance of rinderpest (RP) in accordance with Chapter 1.4. applicable to Members seeking establishment of freedom from RP. Guidance is provided for Members seeking reestablishment of freedom from RP, following an outbreak and for the maintenance of RP free status.

Surveillance strategies employed for demonstrating freedom from RP at an acceptable level of confidence will need to be adapted to the local situation. Outbreaks of rinderpest in cattle may be graded as per-acute, acute or sub-acute. Differing clinical presentations reflect variations in levels of innate host resistance (Bos indicus breeds being more resistant than B. taurus), and variations in the virulence of the attacking strain. Experience has shown that syndromic surveillance strategies i.e. surveillance based on a predefined set of clinical signs (e.g. searching for “stomatitis-enteritis syndrome”) are useful to increase the sensitivity of the system. It is generally accepted that unvaccinated populations of cattle are likely to promote the emergence of virulent strains and associated epidemics while partially vaccinated populations favor the emergence of mild strains associated with endemic situations. In the case of per-acute cases the presenting sign may be sudden death. In the case of sub-acute (mild) cases, clinical signs are irregularly displayed and difficult to detect.

In certain areas there are some key wildlife populations, especially African buffaloes, which act as sentinels for rinderpest infection. These subpopulations should be included in the design of the surveillance strategy.

Surveillance for RP should be in the form of a continuing programme designed to establish that the whole country is free from RP virus (RPV) infection.

Article 8.13.21.

Surveillance: definitions general conditions and methods

1. Rinderpest

For the purpose of this Chapter, rinderpest is defined as an infection of large ruminants (cattle, buffaloes, yaks, etc.), small ruminants, pigs and various wildlife species within the order Artiodactyla, caused by rinderpest virus. In small ruminants and various species of wildlife, particularly antelopes, infection generally passes without the development of frank clinical signs. Characteristic clinical signs and pathological lesions are described in Chapter 2.1.15. of the Terrestrial Manual.
Outbreaks of rinderpest in cattle may be graded as per acute, acute or sub acute. Differing clinical presentations reflect variations in levels of innate host resistance (Bos indicus breeds being more resistant than Bos taurus), and variations in the virulence of the attacking strain. It is generally accepted that unvaccinated populations of cattle are likely to promote the emergence of virulent strains and associated epidemics while partially vaccinated populations favour the emergence of mild strains associated with endemic situations. In the case of per acute cases the presenting sign may be sudden death. In the case of sub acute (mild) cases, clinical signs are irregularly displayed and difficult to detect.

Freedom from rinderpest means freedom from rinderpest virus infection.

1. A surveillance system in accordance with Chapter 1.4. should be under the responsibility of the Veterinary Authority. A procedure should be in place for the rapid collection and transport of samples from suspect cases of RP to a laboratory for RP diagnoses as described in the Terrestrial Manual.

2. Rinderpest vaccines

For the purpose of this Chapter and the Terrestrial Code, OIE-recognised rinderpest vaccines currently in use, or likely to become so in the foreseeable future, are considered to be commercial modified live vaccines produced from attenuated rinderpest virus (referred to as ‘rinderpest vaccine’) produced in accordance with Chapter 2.1.15. of the Terrestrial Manual.

2. The RP surveillance programme should:

   a) include an early warning system throughout the production, marketing and processing chain for reporting suspicious cases. Farmers and workers who have day-to-day contact with livestock, as well as diagnosticians, should report promptly any suspicion of RP. They should be supported directly or indirectly (e.g. through private veterinarians or veterinary para-professionals) by government information programmes and the Veterinary Authority. All significant epidemiological events consistent with “stomatitis-enteritis syndrome” should be investigated immediately. Where suspicion cannot be resolved by epidemiological and clinical investigation, samples should be taken and submitted to a laboratory. This requires that sampling kits and other equipment are available for those responsible for surveillance. Personnel responsible for surveillance should be able to call for assistance from a team with expertise in RP diagnosis and control;

   b) implement, when relevant, regular and frequent clinical inspection and serological testing of high-risk groups of animals, such as those adjacent to an RP infected country.

An effective surveillance system will periodically identify suspicious cases compatible with the “stomatitis-enteritis syndrome” that require follow-up and investigation to confirm or exclude that the cause of the condition is RPV. The rate at which such suspicious cases are likely to occur will differ between epidemiological situations and cannot therefore be predicted reliably. Applications for freedom from RPV infection should, in consequence, provide details of the occurrence of suspicious cases and how they were investigated and dealt with. This should include the results of laboratory testing and the control measures to which the animals concerned were subjected during the investigation (quarantine, movement stand-still orders, etc.).

Article 8.13.22.

Surveillance activities strategies

General recommendations on animal disease surveillance are outlined in Chapter 1.4. of the Terrestrial Code.
Annex XIX (contd)

Rinderpest must be a notifiable disease i.e. notification of outbreaks of rinderpest as soon as detected or suspected must be brought to the attention of the Veterinary Authority.

The precise surveillance information required for establishing freedom will differ from country to country depending on factors such as the former rinderpest status of the country, the regional rinderpest situation and accreditation status, the time elapsing since the last occurrence of rinderpest, livestock husbandry systems (e.g. extensive pastoralism, nomadism and transhumance versus sedentary agropastoralism) and trading patterns.

Evidence of efficiency of the surveillance system can be provided by the use of performance indicators.

Surveillance results presented will be expected to have accrued from a combination of surveillance activities including some or all of the following:

1. A routine national animal disease reporting system supported by evidence of its efficiency and follow-up - an on-going, statutory, centrally organised system of reporting.

Ideally disease reports should be expressed in a Geographical Information System environment and analysed for clustering of observations and followed up.

1. Introduction

The target population for surveillance aimed at identifying disease and infection should cover all significant populations of susceptible species within the country to be recognised as free from RPV infection.

The strategy employed can be based on randomised sampling requiring surveillance consistent with demonstrating the absence of RPV infection at an acceptable level of statistical confidence. The frequency of sampling should be dependent on the epidemiological situation. Targeted surveillance (e.g. based on the increased likelihood of infection in particular localities or species) can be an appropriate strategy. The applicant Member should justify the surveillance strategy chosen as adequate to detect the presence of RPV infection in accordance with Chapter 1.4. and the epidemiological situation. It may, for example, be appropriate to target clinical surveillance at particular subpopulations likely to exhibit clear clinical signs. For targeted surveillance consideration should be given to the following:

i) historical disease patterns (risk mapping) - clinical, participatory and laboratory-based;

ii) critical population size, structure and density;

iii) livestock husbandry and farming systems;

iv) movement and contact patterns - markets and other trade-related movements;

v) transmission parameters (e.g. virulence of the strain, animal movements);

vi) wildlife and other species demography.

For random surveys, the design of the sampling strategy will need to take into account the expected disease prevalence. The sample size selected for testing will need to be large enough to detect infection if it were to occur at a predetermined minimum rate. The sample size and expected disease prevalence determine the level of confidence in the results of the survey. The applicant Member must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation in accordance with Chapter 1.4. Selection of the expected prevalence in particular clearly needs to be based on the prevailing or historical epidemiological situation.
Irrespective of the survey design selected, the sensitivity and specificity of the diagnostic tests employed are key factors in the design, sample size determination and interpretation of the results obtained.

Irrespective of the testing system employed, surveillance design should anticipate the occurrence of false positive reactions. If the characteristics of the testing system are known, the rate at which these false positives are likely to occur can be calculated in advance. There needs to be an effective procedure for following-up positives to ultimately subsequently determine with a high level of confidence whether they are indicative of infection or not. This should involve both supplementary tests and follow-up investigation to collect diagnostic material from the original sampling unit as well as herds which may be epidemiologically linked to it.

The principles involved in surveillance for disease infection are technically well defined in Chapter 1.4. The design of surveillance programmes to prove the absence of RPV infection needs to be carefully followed to ensure the reliability of results. The design of any surveillance programme, therefore, requires inputs from professionals competent and experienced in this field.

2. Emergency disease reporting systems and investigation of epidemiologically significant events ('stomatitis-enteritis syndrome')

Emergency reporting systems can be devised to short-circuit normal passive reporting systems to bring suspicious events to the fore and lead to rapid investigation and tracing. All such investigations should be well documented for presentation as an outcome of the surveillance system.

2. Clinical surveillance

Clinical surveillance aims at detecting clinical signs of “stomatitis-enteritis syndrome” by close physical examination of susceptible animals. Whereas significant emphasis is placed on the diagnostic value of mass serological screening, surveillance based on clinical inspection should not be underrated. It may be able to provide a high level of confidence of detection of disease if sufficiently large numbers of clinically susceptible animals are examined. It is essential that clinical cases detected be followed by the collection of appropriate samples such as ocular and nasal swabs, blood or other tissues for virus isolation. Clinical surveillance and laboratory testing should always be applied in series to clarify the status of RP suspects detected by either of these complementary diagnostic approaches. Laboratory testing may confirm clinical suspicion, while clinical surveillance may contribute to clarification of positive serology. Any sampling unit within which suspicious animals are detected should be classified as infected until contrary evidence is produced.

Active search for clinical disease can include participatory disease searching, tracing backwards and forwards, and follow-up investigations. Participatory disease surveillance is a form of targeted active surveillance based upon methods to capture livestock owners perceptions on the prevalence and patterns of disease.

The often underestimated labour intensity requirements and the logistical difficulties involved in conducting clinical examinations should not be underestimated and should be taken into account.

It is essential that all RPV isolates are sent to an OIE reference laboratory to determine the biological characteristics of the causative virus as well as its genetic and antigenic characterization.
Annex XIX (contd)

3. Detection and thorough investigation of epidemiologically significant events ('stomatitis enteritis syndrome') which raise suspicion of rinderpest supported by evidence of efficiency of the system.

Laboratory examination undertaken to confirm or rule out rinderpest is given extra credibility if it is accompanied by the results of differential diagnostic examinations.

3. Virological surveillance

Given that RP is an acute infection with no known carrier state, virological surveillance using tests described in the Terrestrial Manual should be conducted to confirm clinically suspect cases. Applying virological methods in seropositive animals is not regarded as an efficient approach.

4. Searching for evidence of clinical rinderpest

Active search for disease might include participatory disease searching combined with village disease searching, tracing backwards and forwards, follow-up and investigation.

5. Serosurveillance

Serosurveillance aims at detecting antibodies against RPV. Positive RPV antibody test results can have four possible causes:

a) natural infection with RPV;

b) vaccination against RP;

c) maternal antibodies derived from an immune dam (maternal antibodies in cattle can be found only up to 12 months of age);

d) heterophile (cross) and other non-specific reactions.

a) Randomised serosurveys

Statistically selected samples from relevant strata within the host populations are examined to detect serological evidence of possible virus circulation.

A sampling unit for the purposes of disease investigation and surveillance is defined as a group of animals in sufficiently close contact that individuals within the group are at approximately equal risk of coming in contact with the virus if there should be an infectious animal within the group. In most circumstances, the sampling unit will be a herd which is managed as a unit by an individual or a community, but it may also be other epidemiologically appropriate groupings which are subject to regular mixing, such as all animals belonging to residents of a village. In the areas where nomadic or transhumant movements exist, the sampling unit can be the permanent bore holes, wells or water points. Sampling units should normally be defined so that their size is generally between 50 and 1,000 animals.

i) Criteria for stratification of host populations

Strata are homogeneously mixing sub-populations of livestock. Any disease surveillance activities must be conducted on populations stratified according to the management system, and by herd size where this is variable. Herds, or other sampling units, should be selected by proper random statistical selection procedures from each stratum.
ii) Field procedures and sample sizes

Annual sample sizes shall be sufficient to provide 95% probability of detecting evidence of rinderpest if present at a prevalence of 1% of herds or other sampling units and 5% within herds or other sampling units. This can typically be achieved by examining 300 herds per stratum per year, but procedures for sampling should be in accordance with the “Guide to Epidemiological Surveillance for Rinderpest”, or another procedure that would achieve the same probability of detection.

Where the sampling frame of herds is known, herds shall be selected for examination by the use of random number tables. Otherwise, samples of herds can be selected by taking the nearest herd to a randomly selected map reference, provided that the herds are evenly distributed. Failing this, any herd(s) within a fixed radius of randomly selected map references should be sampled. It must be compulsory for any selected herd to be examined or tested as required.

In carrying out clinical surveillance for evidence of rinderpest, all animals in selected herds or sampling units will be examined by a veterinarian for signs of the disease, especially mouth lesions. Any positive result shall be evaluated using epidemiological and laboratory methods to confirm or refute the suspicion of rinderpest virus activity. All animals born after the cessation of vaccination and more than one year old will be eligible for serological testing.

Where operational considerations require it, the number of eligible animals tested within each sampled herd may be reduced. This will reduce the probability of within herd detection and there must be at least a compensatory increase in the number of herds sampled, so that the required 95% probability of detecting 1% between herd prevalence is maintained.

b) Risk-focused serosurveillance

Risk-focused serosurveillance differs from randomised serosurveillance in that it increases detection sensitivity by obtaining samples from areas/populations determined to be at higher risk of infection, so as to detect serological evidence of possible virus circulation. The operational modalities for risk-based focussing of surveillance require definition (randomisation within defined focus, high risk animals, etc.). The extent to which randomisation needs to be retained in the generation of risk-focused serosurveillance data needs to be established.

Focussing can be achieved by reference to some or all of the following:

i) Historical disease patterns (prior probability mapping) — clinical, participatory and laboratory-based

ii) Critical population size, structure and density

iii) Livestock husbandry and farming systems

iv) Movement and contact patterns — markets and other trade-related movements

v) Transmission parameters (e.g. virulence of the strain, animal movements)

vi) Wildlife and other species demography.
Selection of cattle and buffaloes for serosurveillance

Ageing cattle and Asian buffaloes for the purpose of serosurveillance:

Mis-ageing of cattle selected for serosurveillance is the most common source of error. Colostral immunity can persist almost up to one year of age when measured by the Hc-ELISA. Thus, it is essential to exclude from sampling buffaloes and cattle less than one year of age. In addition, it is frequently necessary to be able to exclude those which are older than a certain age, for example, to select only those born after cessation of vaccination.

Accounts of the ages for eruption of the incisor teeth vary markedly and are clearly dependent on species, breed, nutritional status and nature of the feed.

Pragmatically, and solely for the purposes of serosurveillance, it can be accepted that:

a) cattle having only one pair of erupted permanent central incisor teeth are aged between 21 and 36 months (Asian buffaloes 24-48 months);

b) cattle having only two pairs of erupted permanent central incisor teeth are aged between 30 and 48 months (Asian buffaloes 48-60 months).

Thus selecting a cohort of cattle possessing only one pair of permanent incisors will preclude any interference from maternal immunity derived from earlier vaccination or infection and ensure that vaccinated cattle are not included if vaccination ceased 3 years or more previously (for Asian buffaloes 4 years or more).

It is important to select a cohort of cattle possessing only one pair of permanent incisors to preclude any interference from maternal immunity derived from earlier vaccination or infection and ensure that vaccinated cattle are not included.

Although it is stressed here that animals with milk teeth only are not suitable for surveillance based on serology, they are of particular interest and importance in surveillance for clinical disease. After the loss of colostral immunity, by about one year of age, these are the animals which are most likely to suffer the more severe disease form and in which to look for lesions indicative of rinderpest.

It may be possible to use serum collected for other survey purposes for RP surveillance. However, the principles of survey design described in this Chapter and the requirement for a statistically valid survey for the presence of RPV should not be compromised.

The discovery of clustering of seropositive reactions should be foreseen. It may reflect any of a series of events, including but not limited to the demographics of the population sampled, vaccinal exposure or the presence of field strain infection. As clustering may signal field strain infection, the investigation of all instances must be incorporated in the survey design.

The results of random or targeted serological surveys are important in providing reliable evidence that RPV infection is not present in a country. It is therefore essential that the survey be adequately thoroughly documented.
Article 8.13.24.

Wildlife surveillance where a significant susceptible wildlife population exists

There are some key wildlife populations, especially African buffaloes, which act as sentinels for rinderpest infection. Where a significant population of a susceptible wildlife species exists, serosurveillance data are required to support absence of infection. These populations should be monitored purposively to support the dossiers to be submitted for freedom from rinderpest virus infection. Detection of virus circulation in wildlife can be undertaken indirectly by sampling contiguous livestock populations.

Obtaining meaningful data from wildlife surveillance can be enhanced by close coordination of activities in the regions and countries. Both purposive and opportunistic samplings are used to obtain material for analysis in national and reference laboratories. The latter are required because most many countries are unable do not have adequate facilities to perform the full testing protocol for detecting rinderpest RP antibodies in wildlife sera.

Purposive Targeted sampling is the preferred method to provide wildlife data to evaluate the status of rinderpest infection. In reality, the capacity to perform purposive work targeted surveillance in the majority of countries remains minimal. Opportunistic sampling (hunting) is feasible and it provides However, samples can be obtained from hunted animals, and these may provide useful background information.

Wildlife form transboundary populations; therefore, any data from the population could be used to represent the result for the ecosystem and be submitted by more than one Member in an application to the OIE (even if the sampling was not obtained in the territory of the OIE Member submitting the application). It is therefore recommended therefore that the Countries or Territories represented in a particular ecosystem should coordinate their sampling programmes.

The standards for serosurveillance are different from that set for cattle because the serological tests are not fully validated for wildlife species and financial and logistic constraints of sampling prevent collection of large numbers of samples.

Where the serological history of the herd is known from previous work (as might be the case for a sentinel herd), repeat sampling need only focus on the untested age groups, born since the last known infection. The sample needs to be taken according to the known epidemiology of the disease in a given species. Opportunistic Samples collected from hunted animals, which are positive, should not be interpreted without a targeted survey to confirm the validity of these results. Opportunistic Such sampling cannot follow a defined protocol and therefore can only provide background information.

From the collective experience of the laboratories and experts over the years, an appropriate test protocol is based on the high expected sero-prevalence in a previously infected buffalo herd (99% seroconversion of eligible animals within a herd), which is detected using a test, which is 100% sensitive. No single test can achieve this, however, combining H o-ELISA to VNT raises sensitivity close to 100%.

In the order of 1-2% of a herd of African buffaloes must be sampled to ensure that no positive case is missed. For example in a herd of 300 buffaloes, five animals should be sampled and the above multiple test protocol followed. Where the serological history of the herd is known from previous work (as might be the case for a sentinel herd), repeat sampling need only focus on the untested age groups, born since the last known infection. Appropriate sampling fraction for other wildlife species are less well defined, as social organization (herd structure, likely contact rates, etc.) vary. The sample needs to be taken according to the known epidemiology of the disease in a given species. Opportunistic sampling which are positive, should not be interpreted without a purposive survey to confirm the validity of these results. Opportunistic sampling cannot follow a defined protocol and therefore can only provide background information.
Annex XIX (contd)

Article 8.13.25.

**Evaluation of applications for accreditation of Members applying for recognition of freedom from rinderpest (RP)**

Evaluation of applications for the status of freedom from rinderpest will be the responsibility of the OIE Scientific Commission for Animal Diseases which can request the Director General of the OIE to appoint an ad hoc group in order to assist in reaching an informed decision to present to the OIE International Committee for approval.

The composition and method of selection of the ad hoc group shall be such as to ensure both a high level of expertise in evaluating the evidence and total independence of the group in reaching conclusions concerning the disease status of a particular country.

In addition to the general conditions described in this Chapter, a Member applying for recognition of RP freedom for the country should provide evidence for the existence of an effective surveillance programme. The strategy and design of the surveillance programme will depend on the prevailing epidemiological circumstances and will be planned and implemented according to general conditions and methods in this Chapter, to demonstrate absence of RPV infection, during the preceding 24 months in susceptible populations. This requires the support of a national or other laboratory able to undertake identification of RPV infection through virus/antigen/genome detection and antibody tests described in the Terrestrial Manual.


**Steps to be taken to declare a country to be free from rinderpest**

Recognition of the status ‘free from rinderpest’ is given to a Member. Where traditionally managed livestock move freely across international borders, groups of Members may usefully associate themselves into a group for the purposes of obtaining data to be used for mutually supportive applications for individual country accreditation.

For the purpose of this Chapter, the following assumptions are made:

a) that within most previously infected countries, rinderpest vaccine will have been used to control the rate of infection;

b) that within an endemically infected population there will be a large number of immune hosts (both vaccinates and recovered animals);

c) that the presence of a proportion of immune hosts within a vaccinated population could have led to a slowing of the rate of virus transmission and possibly the concomitant emergence of strains of reduced virulence, difficult to detect clinically;

d) that the virulence of the virus (and therefore the ease of clinical detection) may or may not increase as the herd immunity declines following withdrawal of vaccination; however, continuing transmission will generate serological evidence of their persistence.
Before accreditation can be considered, countries which have the disease by the use of rinderpest vaccine must wait until an unvaccinated cohort is available to allow meaningful serological surveillance to be conducted.

The OIE has concluded that the majority of countries have stopped vaccinating for a sufficient length of time for it now to be feasible that a single submission of evidence gained over 2 years of appropriate surveillance shall be sufficient to gain rinderpest free accreditation.

A Member accredited as free from rinderpest must thereafter submit annual statements to the Director General of the OIE indicating that surveillance has failed to disclose the presence of rinderpest, and that all other criteria continue to be met.

A country previously infected with rinderpest which has not employed rinderpest vaccine for at least 25 years and has throughout that period detected no evidence of rinderpest virus disease or infection may be accredited as free from rinderpest by the OIE based on historical grounds, provided that the country:

- has had throughout at least the last 10 years and maintains permanently an adequate animal disease surveillance system along with the other requirements outlined in Article 8.13.25;

- is in compliance with OIE reporting obligations (Chapter 1.1.).

The Veterinary Authorities of the Member must submit a dossier containing evidence supporting their claim to be free from rinderpest on a historical basis to the Director General of the OIE for evaluation by the OIE Scientific Commission for Animal Diseases and accreditation by the OIE International Committee. The dossier should contain at least the following information:

- a description of livestock populations, including wildlife;

- the history of rinderpest occurrence in the country and its control;

- an affirmation that rinderpest has not occurred for 25 years, that vaccine has not been used during that time, and that rinderpest is a notifiable disease;

- evidence that in the last 10 years the disease situation throughout the Member has been constantly monitored by a competent and effective veterinary infrastructure that has operated a national animal disease reporting system submitting regular (monthly) disease occurrence reports to the Veterinary Authority;

- the structure and functioning of the Veterinary Services;

- the Member operates a reliable system of risk analysis based importation of livestock and livestock products.

Evidence in support of these criteria must accompany the Member's accreditation application dossier. In the event that satisfactory evidence is not forthcoming, the OIE may seek clarification or refer the dossier back to the originators, giving its reasons for so doing. Under such circumstances a fresh dossier would be entertained in due course.
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A Member having eradicated rinderpest within the last 25 years, wishing to be accredited free from rinderpest and having ended rinderpest vaccination must initiate a two-year surveillance programme to demonstrate freedom from rinderpest whilst banning further use of rinderpest vaccine. The step of accreditation as free from rinderpest is subject to meeting stringent criteria with international verification under the auspices of the OIE.

A country historically infected with rinderpest but which has convincing evidence that the disease has been excluded for at least 2 years and is not likely to return, may apply to OIE to be accredited as free from rinderpest. The conditions which apply include that an adequate animal disease surveillance system has been maintained throughout at least that period.

The Veterinary Authority of the Member must submit a dossier containing evidence supporting their claim to be free from rinderpest to the Director General of the OIE for evaluation by the OIE Scientific Commission for Animal Diseases and accreditation by the OIE International Committee showing that they comply with:

- the provisions outlined in this Chapter;
- OIE reporting obligations outlined in Chapter 1.1. of the Terrestrial Code.

Other conditions that apply are:

- The Member affirms that rinderpest has not occurred for at least 2 years, that vaccine has not been used during that time, and that rinderpest is a notifiable disease.
- The Veterinary Authority has issued orders curtailing the distribution and use of rinderpest vaccine in livestock.
- The Veterinary Authority has issued orders for the recall and destruction of rinderpest vaccine already issued.
- The Veterinary Authority has issued orders restricting the importation of rinderpest vaccine into, or the further manufacture of rinderpest vaccine within, the territory under his jurisdiction. An exception can be made for establishing a safeguarded rinderpest emergency vaccine bank under the control of the Chief Veterinary Officer who can demonstrate that no calls have been made on that vaccine bank.
- The Veterinary Authority has set in place a rinderpest contingency plan.
- Over the previous 2 years at least, the disease situation throughout the Member has been constantly monitored by a competent and effective infrastructure that has operated a national animal disease reporting system submitting regular (monthly) disease occurrence reports to the Veterinary Authority.
- All outbreaks of disease with a clinical resemblance to rinderpest have been thoroughly investigated and routinely subjected to laboratory testing by an OIE recognised rinderpest-specific test within the national rinderpest laboratory or at a recognised reference laboratory.
Annex XIX (contd)

The dossier shall contain:

- the results of a continuous surveillance programme, including appropriate serological surveys conducted during at least the last 24 months, providing convincing evidence for the absence of rinderpest-virus circulation;
- a description of livestock populations including wildlife;
- the history of rinderpest occurrence in the country and its control;
- an affirmation that rinderpest has not occurred for at least 2 years, that vaccine has not been used during that time, and that rinderpest is a notifiable disease;
- evidence that in the last 2 years the disease situation throughout the Member has been constantly monitored by a competent and effective veterinary infrastructure that has operated a national animal disease reporting system submitting regular (monthly) disease occurrence reports to the Veterinary Authority;
- the structure and functioning of the Veterinary Services;
- the Member operates a reliable system of risk analysis based importation of livestock and livestock products.

In the event that satisfactory evidence in support of the application is not forthcoming, the OIE may seek clarification or refer the dossier back to the originators, giving its reasons for so doing. Under such circumstances a fresh dossier would be entertained in due course.

Article 8.13.2726.

Rinderpest outbreaks after accreditation and recovery of rinderpest-free status Members re-applying for recognition of freedom from RP following an outbreak

Should there be an outbreak, or outbreaks, of rinderpest in a Member at any time after recognition of rinderpest freedom, the origin of the virus strain must be thoroughly investigated. In particular it is important to determine if this is due to the re-introduction of virus or re-emergence from an undetected focus of infection. The virus must be isolated and compared with historical strains from the same area as well as those representatives of other possible sources. The outbreak itself must be contained with the utmost rapidity using the resources and methods outlined in the Contingency Plan.

Following an outbreak, or outbreaks, of rinderpest in a Member at any time after recognition of rinderpest freedom, the origin of the virus strain should be thoroughly investigated. In particular it is important to determine if this is due to the re-introduction of virus or re-emergence from an undetected focus of infection. Ideally, the virus should be isolated and compared with historical strains from the same area as well as those representatives of other possible sources.

After elimination of the outbreak, a Member wishing to regain the status ‘free from rinderpest’ must undertake serosurveillance according to this Chapter to determine the extent of virus spread. In addition to the general conditions described in this Chapter, a Member re-applying for recognition of country freedom from RP should show evidence of an active surveillance programme for RP as well as absence of RPV infection.
If investigations show the outbreak virus originated from outside the country, provided the outbreak was localised, rapidly contained and speedily eliminated, and provided there was no serological evidence of virus spread outside the index infected area, accreditation of freedom could proceed rapidly. The Member must satisfy the OIE Scientific Commission for Animal Diseases that the outbreaks were contained, eliminated and did not represent endemic infection.

An application to regain the status free from rinderpest shall not generally be accepted until both clinical and serological evidence shows that there has been no virus transmission for at least 3 or 6 months, depending on whether or not stamping out or vaccination respectively has been applied.

Article 8.3.27.

The use and interpretation of serological tests for serosurveillance of RP

Serological testing is an appropriate tool to use for RP surveillance. The prescribed serological tests which should be used for RP surveillance are described in the Terrestrial Manual; these are of high diagnostic specificity and minimise the proportion of false positive reactions. Antibodies to virulent strains and the Kabete O vaccine strain of RPV can be detected in cattle from about 10 days post infection (approximately 7 days after the appearance of fever) and peak around 30 to 40 days post infection. Antibodies then persist for many years, possibly for life, although titres decline with time. In the case of less virulent strains the detection of the antibody response by ELISA may be delayed by as much as three weeks. There is only one serotype of virus and the tests will detect antibodies elicited by infection with all RP viruses but the tests cannot discriminate between antibodies to field infection and those from vaccination with attenuated vaccines. This fact compromises serosurveillance in vaccinated populations and realistically meaningful serosurveillance can only commence once vaccination has ceased for several years. In these circumstances, dental ageing of cattle and buffaloes is of great value to minimise the inclusion of animals seropositive by virtue of colostral immunity and historic vaccination or infection. The cohort of cattle with one single set of central incisors is the most appropriate to sample.

The test most amenable to the mass testing of sera as required to demonstrate freedom from infection is the H c-ELISA. Practical experience from well-controlled serological surveillance in non-vaccinated populations in Africa and Asia demonstrate that one can expect false positive reactions in 0.05 % or less of sera tested. The sensitivity of the test approaches 100 % (relative to the VNT) in Kabete O vaccinated cattle and infection with highly virulent viruses but is lower in the case of low virulence strains. Experience supported by experimental studies indicates that in all cases sensitivity exceeds 70 %.

Only tests approved by OIE as indicated in the Terrestrial Manual should be used to generate data presented in support of applications for accreditation of RP freedom. It is necessary to demonstrate that apparently positive serological results have been adequately investigated. The follow-up studies should use appropriate clinical, epidemiological, serological and virological investigations. By this means the investigation should examine all evidence that might confirm or refute the hypothesis that the positive results to the serological tests employed in the survey were not due to virus circulation.
The prescribed serological tests have not been fully validated for use in all wild species. From the collective experience of the reference laboratories and experts over the years, an appropriate test protocol for wildlife is based on the high expected sero-prevalence in a previously infected buffalo herd which is 99% seroconversion of eligible animals within a herd as detected by use of a 100% sensitive test. No single test can achieve this but combining the Hc-ELISA with the VNT raises sensitivity close to 100%.


2. Pragmatically and solely for the purposes of serosurveillance, it can be accepted that:
   a) Cattle having one pair of erupted permanent central incisor teeth are aged between 21 and 36 months (Asian buffaloes 24 to 48 months).
   b) Cattle having only two pairs of erupted permanent central incisor teeth are aged between 30 and 48 months (Asian buffaloes 48-60 months).
CHAPTER 9.1.

ACARAPISOSIS OF HONEY BEES

Article 9.1.1.

General provisions

For the purposes of this Chapter, acarapisosis, acarine disease or tracheal mite infestation is a disease of the adult honey bee *Apis mellifera* L., and possibly of other *A* pis species (such as *A* pis *ɔrana*). It is caused by the Tarsonemid mite *Acarapis woodi* (Rennie). The mite is an internal obligate parasite of the respiratory system, living and reproducing mainly in the large prothoracic trachea of the bee. Early signs of infection normally go unnoticed, and only when infection is heavy does it become apparent; this is generally in the early spring. The infection spreads by direct contact from adult bee to adult bee, with newly emerged bees under 10 days old being the most susceptible. The mortality rate may range from moderate to high.

Standards for diagnostic tests are described in the *Terrestrial Manual*.

Article 9.1.1.bis

Trade in commodities

When authorising import or transit of the following commodities, *Veterinary Authorities* should not require any acarapisosis related conditions, regardless of the acarapisosis status of the honey bee population of the exporting country or zone:

1. honey bee semen and honey bee venom;
2. used equipment associated with beekeeping;
3. honey, beeswax, honey bee-collected pollen, propolis and royal jelly.

When authorising import or transit of other commodities listed in this Chapter, *Veterinary Authorities* should require the conditions prescribed in this Chapter relevant to the acarapisosis status of the honey bee population of the exporting country or zone.

Article 9.1.2.

Determination of the acarapisosis status of a country or zone/compartment

The acarapisosis status of a country or *zone/compartment* (under study) can only be determined after considering the following criteria:

1. a risk assessment has been conducted, identifying all potential factors for acarapisosis occurrence and their historic perspective;
2. acarapisosis should be notifiable in the whole country or *zone/compartment* (under study) and all clinical signs suggestive of acarapisosis should be subjected to field and laboratory investigations;
Annex XX (contd)

3. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of acarapisosis;

4. the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees should have current knowledge of, and authority over, all domesticated apiaries in the whole country.

Article 9.1.3.

Country or zone/compartment (under study) free from acarapisosis

1. Historically free status

A country or zone/compartment (under study) may be considered free from acarapisosis after conducting a risk assessment as referred to in Article 9.1.2. but without formally applying a specific surveillance programme if the country or zone/compartment (under study) complies with the provisions of Chapter 1.4.

2. Free status as a result of an eradication programme

A country or zone/compartment (under study) which does not meet the conditions of point 1 above may be considered free from acarapisosis after conducting a risk assessment as referred to in Article 9.1.2. and when:

a) the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees has current knowledge of, and authority over, all domesticated apiaries existing in the country or zone/compartment (under study);

b) acarapisosis is notifiable in the whole country or zone/compartment (under study), and any clinical cases suggestive of acarapisosis are subjected to field and laboratory investigations;

c) for the 3 years following the last reported case of acarapisosis, annual surveys supervised by the Veterinary Authority, with negative results, have been carried out on a representative sample of apiaries in the country or zone/compartment (under study) to provide a confidence level of at least 95% of detecting acarapisosis if at least 1% of the apiaries were infected at a within-apiary prevalence rate of at least 5% of the hives; such surveys may be targeted towards apiaries, areas and seasons with a higher likelihood of disease;

d) to maintain free status, an annual survey supervised by the Veterinary Authority, with negative results, is carried out on a representative sample of apiaries in the country or zone/compartment (under study) to indicate that there has been no new cases; such surveys may be targeted towards areas with a higher likelihood of disease;

e) (under study) there is no self-sustaining feral population of A. mellifera or other possible host species in the country or zone/compartment (under study);

f) the importation of the commodities listed in this Chapter into the country or zone/compartment (under study) is carried out in conformity with the recommendations of this Chapter.
Recommendations on safe commodities

Regardless of the acarapisosis status of the exporting country, Veterinary Authorities should authorise without restriction the import or transit through their territory of the following commodities:

1. honey bee semen and honey bee venom;
2. used equipment associated with beekeeping;
3. honey, beeswax, honey bee-collected pollen, propolis and royal jelly.

Article 9.1.5.

Recommendations for the importation of live queen honey bees, worker bees and drones with or without associated brood combs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the bees come from a country or zone/compartment (under study) free from acarapisosis.

Article 9.1.6.

Recommendations for the importation of eggs, larvae and pupae of honey bees

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. were sourced from an officially free country or zone/compartment (under study); or
2. were examined by an official laboratory and declared free of all life stages of A. woodi; or
3. have originated from queens in a quarantine station and were examined microscopically and found free of all life stages of A. woodi.

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CHAPTER 9.2.

AMERICAN FOULBROOD OF HONEY BEES

Article 9.2.1.

General provisions

For the purposes of this Chapter, American foulbrood is a disease of the larval and pupal stages of the honey bee *Apis mellifera* and other *Apis* spp., and occurs in most countries where such bees are kept. *Paenibacillus larvae* subsp. *larvae*, the causative organism, is a bacterium that can produce over one billion spores in each infected larva. The spores are very long-living and extremely resistant to heat and chemical agents, and only the spores are capable of inducing the disease.

Combs of infected apiaries may show distinctive clinical signs which can allow the disease to be diagnosed in the field. However, subclinical infections are common and require laboratory diagnosis.

For the purposes of the Terrestrial Code, the incubation period for American foulbrood shall be 15 days (not including the wintering period which may vary according to country).

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 9.2.1.bis

Trade in commodities

When authorising import or transit of the following commodities, Veterinary Authorities should not require any American foulbrood related conditions, regardless of the American foulbrood status of the honey bee population of the exporting country or zone:

1. honey bee semen;
2. honey bee venom.

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the American foulbrood status of the honey bee population of the exporting country or zone.

Article 9.2.2.

Determination of the American foulbrood status of a country or zone/compartment

The American foulbrood status of a country or zone/compartment (under study) can only be determined after considering the following criteria:

1. a risk assessment has been conducted, identifying all potential factors for American foulbrood occurrence and their historic perspective;
2. American foulbrood should be notifiable in the whole country or zone/compartment (under study) and all clinical signs suggestive of American foulbrood should be subjected to field and/or laboratory investigations;
Annex XX (contd)

3. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of American foulbrood;

4. the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of disease of honey bees should have current knowledge of, and authority over, all domesticated apiaries in the country.

Article 9.2.3.

Country or zone/compartment (under study) free from American foulbrood

1. Historically free status

A country or zone/compartment (under study) may be considered free from the disease after conducting a risk assessment as referred to in Article 9.2.2. but without formally applying a specific surveillance programme if the country or zone/compartment (under study) complies with the provisions of Chapter 1.4.

2. Free status as a result of an eradication programme

A country or zone/compartment (under study) which does not meet the conditions of point 1 above may be considered free from American foulbrood after conducting a risk assessment as referred to in Article 9.2.2. and when:

a) the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of disease of honey bees has current knowledge of, and authority over, all domesticated apiaries existing in the country or zone/compartment (under study);

b) American foulbrood is notifiable in the whole country or zone/compartment (under study), and any clinical cases suggestive of American foulbrood are subjected to field and/or laboratory investigations;

c) for the 5 years following the last reported isolation of the American foulbrood agent, annual surveys supervised by the Veterinary Authority, with negative results, have been carried out on a representative sample of apiaries in the country or zone/compartment (under study) to provide a confidence level of at least 95% of detecting American foulbrood if at least 1% of the apiaries were infected at a within-apiary prevalence rate of at least 5% of the hives; such surveys may be targeted towards areas with the last reported isolation of the American foulbrood agent;

d) to maintain free status, an annual survey supervised by the Veterinary Authority, with negative results, is carried out on a representative sample of hives in the country or zone/compartment (under study) to indicate that there has been no new isolations; such surveys may be targeted towards areas with a higher likelihood of isolation;

e) (under study) there is no self-sustaining feral population of A. mellifera or other possible host species in the country or zone/compartment (under study);

f) all equipment associated with previously infected apiaries has been sterilised or destroyed;

g) the importation of the commodities listed in this Chapter into the country or zone/compartment (under study) is carried out in conformity with the recommendations of this Chapter.
Article 9.2.4.

Recommendations on safe commodities

Regardless of the American foulbrood status of the exporting country, Veterinary Authorities should authorise without restriction the import or transit through their territory of honey bee semen and honey bee venom.

Article 9.2.5.

Recommendations for the importation of live queen honey bees, worker bees and drones with or without associated brood combs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the bees come from a country or zone/compartment (under study) officially free from American foulbrood.

Article 9.2.6.

Recommendations for the importation of eggs, larvae and pupae of honey bees

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. were sourced from a free country or zone/compartment (under study); or
2. have been isolated from queens in a quarantine station and all workers which accompanied the queen or a representative sample of eggs or larvae were examined for the presence of P. larvae by bacterial culture or PCR in accordance with the Terrestrial Manual.

Article 9.2.7.

Recommendations for the importation of used equipment associated with beekeeping

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the equipment was sterilised under the supervision of the Veterinary Authority by either immersion in 1% sodium hypochlorite for at least 30 minutes (suitable only for non-porous materials such as plastic and metal), gamma irradiation using a cobalt-60 source at a dose rate of 10 kGy, or processing to ensure the destruction of both bacillary and spore forms of P. larvae in conformity with one of the procedures referred to in Chapter X.X. (under study).

Article 9.2.8.

Recommendations for the importation of honey, honey bee-collected pollen, beeswax, propolis and royal jelly

Veterinary Authorities of importing countries officially free from American foulbrood should require the presentation of an international veterinary certificate attesting that the products:

1. were collected in a country or zone/compartment (under study) free from American foulbrood; or
2. have been processed to ensure the destruction of both bacillary and spore forms of P. larvae in conformity with one of the procedures referred to in Chapter X.X. (under study).

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CHAPTER 9.3.

EUROPEAN FOULBROOD OF HONEY BEES

Article 9.3.1.

General provisions

For the purposes of this Chapter, European foulbrood is a disease of the larval and pupal stages of the honey bee Apis mellifera and other Apis spp., and occurs in most countries where such bees are kept. The causative agent is the non-sporeulating bacterium Melissococcus pluton plutonius. Subclinical infections are common and require laboratory diagnosis. Infection remains enzootic because of mechanical contamination of the honeycombs. Recurrences of disease can therefore be expected in subsequent years.

For the purposes of the Terrestrial Code, the incubation period for European foulbrood shall be 15 days (not including the wintering period which may vary according to country).

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 9.3.1.bis

Trade in commodities

When authorising import or transit of the following commodities, Veterinary Authorities should not require any European foulbrood related conditions, regardless of the European foulbrood status of the honey bee population of the exporting country or zone:

1. honey bee semen;
2. honey bee venom.

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the European foulbrood status of the honey bee population of the exporting country or zone.

Article 9.3.2.

Determination of the European foulbrood status of a country or zone/compartment

The European foulbrood status of a country or zone/compartment (under study) can only be determined after considering the following criteria:

1. a risk assessment has been conducted, identifying all potential factors for European foulbrood occurrence and their historic perspective;
2. European foulbrood should be notifiable in the whole country or zone/compartment (under study) and all clinical signs suggestive of European foulbrood should be subjected to field and laboratory investigations;
Annex XX (contd)

3. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of European foulbrood;

4. the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees should have current knowledge of, and authority over, all apiaries in the whole country.

Article 9.3.3.

Country or zone/compartment (under study) free from European foulbrood

1. Historically free status

A country or zone/compartment (under study) may be considered free from the disease after conducting a risk assessment as referred to in Article 9.3.2. but without formally applying a specific surveillance programme if the country or zone/compartment (under study) complies with the provisions of Chapter 1.4.

2. Free status as a result of an eradication programme

A country or zone/compartment (under study) which does not meet the conditions of point 1 above may be considered free from European foulbrood after conducting a risk assessment as referred to in Article 9.3.2. and when:

a) the Veterinary Authority or other Competent Authority with responsibility for the health, reporting and control of diseases of honey bees has current knowledge of, and authority over, all domesticated apiaries existing in the country or zone/compartment (under study);

b) European foulbrood is notifiable in the whole country or zone/compartment (under study), and any clinical cases suggestive of European foulbrood are subjected to field and laboratory investigations;

c) for the 3 years following the last reported isolation of the European foulbrood agent, an annual survey supervised by the Veterinary Authority, with negative results, have been carried out on a representative sample of apiaries in the country or zone/compartment (under study) to provide a confidence level of at least 95% of detecting European foulbrood if at least 1% of the apiaries were infected at a within-apiary prevalence rate of at least 5% of the hives; such surveys may be targeted towards areas with the last reported isolation of the European foulbrood agent;

d) to maintain free status, an annual survey supervised by the Veterinary Authority, with negative results, is carried out on a representative sample of hives in the country or zone/compartment (under study) to indicate that there has been no new isolations; such surveys may be targeted towards areas with a higher likelihood of isolation;

e) (under study) there is no self-sustaining feral population of A. mellifera or other possible host species in the country or zone/compartment (under study);

f) the importation of the commodities listed in this Chapter into the country or zone/compartment (under study) is carried out in conformity with the recommendations of this Chapter.
Article 9.3.4.

Recommendations on safe commodities

Regardless of the European foulbrood status of the exporting country, Veterinary Authorities should authorise without restriction the import or transit through their territory of honey bee semen and honey bee venom.

Article 9.3.5.

Recommendations for the importation of live queen honey bees, worker bees and drones with or without associated brood combs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the bees come from a country or zone/compartment (under study) free from European foulbrood.

Article 9.3.6.

Recommendations for the importation of eggs, larvae and pupae of honey bees

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. were sourced from a free country or zone/compartment (under study); or

2. have been isolated from queens in a quarantine station, and all workers which accompanied the queen or a representative sample of eggs or larvae were examined for the presence of **Melissococcus pluton** by bacterial culture or PCR in accordance with the Terrestrial Manual.

Article 9.3.7.

Recommendations for the importation of used equipment associated with beekeeping

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the equipment was sterilised under the supervision of the Veterinary Authority by either immersion in 0.5% sodium hypochlorite for at least 20 minutes (suitable only for non-porous materials such as plastic and metal), gamma irradiation using a cobalt-60 source at a dose rate of 10 kGy, or processing to ensure the destruction of **Melissococcus pluton**, in conformity with one of the procedures referred to in Chapter X.X. (under study).

Article 9.3.8.

Recommendations for the importation of honey, honey bee-collected pollen, beeswax, propolis and royal jelly

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:
Annex XX (contd)

1. were collected in a country or zone/ compartment (under study) free from European foulbrood; or

2. have been processed to ensure the destruction of *Melissococcus pluton* *M. plutonius* in conformity with one of the procedures referred to in Chapter X.X. (under study).

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CHAPTER 9.4.

SMALL HIVE BEETLE INFESTATION
(Aethina tumida)

Article 9.4.1.

General provisions

For the purposes of this Chapter, small hive beetle (SHB) is an infestation of bee colonies by the beetle Aethina tumida, which is a free-living predator and scavenger affecting populations of the honey bee Apis mellifera L. It can also parasitise bumble bee Bombus terrestris colonies under experimental conditions, and although infestation has not been demonstrated in wild populations, Bombus spp. must also be considered to be susceptible to infestation.

The adult beetle is attracted to bee colonies to reproduce, although it can survive and reproduce independently in other natural environments, using other food sources, including certain types of fruit. Hence once it is established within a localised environment, it is extremely difficult to eradicate.

The life cycle of A. tumida begins with the adult beetle laying eggs within infested hives. These are usually laid in irregular masses in crevices or brood combs. After 2-6 days, the eggs hatch and the emerging larvae begin to feed voraciously on brood comb, bee eggs, pollen and honey within the hive. The SHB has a high reproductive potential. Each female can produce about 1,000 eggs in its 4 to 6 months of life. At maturation (approximately 10-29 days after hatching), the larvae exit the hive and burrow into soil around the hive entrance. Adult beetles emerge after an average of 3-4 weeks, although pupation can take between 8 and 60 days depending on temperature and moisture levels.

The life span of an adult beetle depends on environmental conditions such as temperature and humidity but, in practice, adult beetles can live for at least 6 months and, in favourable reproductive conditions, the female is capable of laying new egg batches every 5-12 weeks. The beetle is able to survive at least 2 weeks without food and 50 days on brood combs.

Early signs of infestation may go unnoticed, but the growth in the beetle population is rapid, leading to high bee mortality in the hive. Because A. tumida can be found and can thrive within the natural environment, and can fly up to 6-13 km from its nest site, it is capable of dispersing rapidly and directly colonising hives. Dispersal includes following or accompanying swarms. Spread of infestation does not require contact between adult bees. However, the movement of adult bees, honeycomb and other apiculture products and used equipment associated with bee-keeping may all cause infestations to spread to previously unaffected colonies.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 9.4.1.bis

Trade in commodities

When authorising import or transit of the following commodities, Veterinary Authorities should not require any small hive beetle infestation related conditions, regardless of the A. tumida status of the honey bee and bumble bee population of the exporting country or zone.
Annex XX (contd)

1. honey bee semen and honey bee venom;
2. packaged extracted honey, refined or rendered beeswax, propolis and frozen or dried royal jelly.

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the A. tumida status of the honey bee and bumble bee population of the exporting country or zone.

Article 9.4.2.

Determination of the A. tumida status of a country or zone

The A. tumida status of a country or zone can only be determined after considering the following criteria:

1. A. tumida infestation should be notifiable in the whole country, and all signs suggestive of A. tumida infestation should be subjected to field and laboratory investigations;
2. on-going awareness and training programmes should be in place to encourage reporting of all cases suggestive of A. tumida infestation;
3. the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees should have current knowledge of, and authority over, all domesticated apianes in the country.

Article 9.4.3.

Country or zone free from A. tumida

1. Historically free status

A country or zone may be considered free from the pest after conducting a risk assessment as referred to in Article 9.4.2. but without formally applying a specific surveillance programme if the country or zone complies with the provisions of Chapter 1.4.

2. Free status as a result of an eradication programme

A country or zone which does not meet the conditions of point 1 above may be considered free from A. tumida infestation after conducting a risk assessment as referred to in Article 9.4.2. and when:

a) the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees has current knowledge of, and authority over, all domesticated apianes existing in the country or zone;

b) A. tumida infestation is notifiable in the whole country or zone, and any clinical cases suggestive of A. tumida infestation are subjected to field and laboratory investigations; a contingency plan is in place describing controls and inspection activities;
c) for the 5 years following the last reported case of A. tumida infestation, an annual survey supervised by the Competent Veterinary Authority, with negative results, has been carried out on a representative sample of apiaries in the country or zone to provide a confidence level of at least 95% of detecting A. tumida infestation if at least 1% of the apiaries were infested at a within-apiary prevalence rate of at least 5% of the hives; such surveys may be targeted towards areas with a higher likelihood of infestation;

d) to maintain free status, an annual survey supervised by the Competent Veterinary Authority, with negative results, is carried out on a representative sample of apiaries to indicate that there have been no new cases; such surveys may be targeted towards areas with a higher likelihood of infestation;

e) all equipment associated with previously infested apiaries has been destroyed, or cleaned and sterilised to ensure the destruction of A. tumida spp., in conformity with one of the procedures referred to in Chapter X.X. (under study);

f) the soil and undergrowth in the immediate vicinity of all infested apiaries has been treated with a soil drench or similar suitable treatment that is efficacious in destroying incubating A. tumida larvae and pupae;

g) the importation of the commodities listed in this Chapter into the country or zone is carried out, in conformity with the recommendations of this Chapter.

Article 9.4.4.

Recommendations on safe commodities

Regardless of the status of the exporting country with regard to A. tumida infestation, Competent Veterinary Authorities should authorise without restriction the import or transit through their territory of the following commodities:

1. honey bee semen and honey bee venom;

2. packaged extracted honey, refined or rendered beeswax, propolis and frozen or dried royal jelly.

Article 9.4.5.

Recommendations for the importation of individual consignments containing a single live queen honey bee or queen bumble bee, accompanied by a small number of associated attendants (a maximum of 20 attendants per queen)

Competent Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the bees come from a country or zone officially free from A. tumida infestation.

OR

Competent Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate including an attestation from the Competent Veterinary Authority of the exporting third country stating that:
Annex XX (contd)

1. the bees come from hives or colonies which were inspected immediately prior to dispatch and show no signs or suspicion of the presence of A. tumida or its eggs, larvae or pupae; and

2. the bees come from an area of at least 100 km radius where no apiary has been subject to any restrictions associated with the occurrence of A. tumida for the previous 6 months; and

3. the bees and accompanying packaging presented for export have been thoroughly and individually inspected and do not contain A. tumida or its eggs, larvae or pupae; and

4. the consignment of bees is covered with fine mesh through which a live beetle cannot enter.

Article 9.4.6.

Recommendations for the importation of live worker bees, drone bees or bee colonies with or without associated brood combs or for live bumble bees

Competent Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the bees come from a country or zone officially free from A. tumida infestation; and

2. the bees and accompanying packaging presented for export have been inspected and do not contain A. tumida or its eggs, larvae or pupae; and

3. the consignment of bees is covered with fine mesh through which a live beetle cannot enter.

Article 9.4.7.

Recommendations for the importation of eggs, larvae and pupae of honey bees or bumble bees

Competent Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the products were sourced from a free country or zone free from A. tumida infestation (under study); OR

2. have been isolated from queens in a quarantine station; and

2. the products have been bred and kept under a controlled environment within a recognised establishment which is supervised and controlled by the Veterinary Authority;

3. are from hives or come from hives or colonies which were inspected immediately prior to entry into the quarantine station and show no signs or suspicion of the presence of A. tumida or its eggs or larvae or pupae then and during the quarantine period.

3. the establishment referred to above was inspected immediately prior to dispatch and all eggs, larvae and pupae show no clinical signs or suspicion of the presence of A. tumida or its eggs or larvae or pupae and
Annex XX (cont’d)

4. the packaging material, containers, accompanying products and food are new and all precautions have been taken to prevent contamination with A. tumida or its eggs, larvae or pupae.

Article 9.4.8.

Recommendations for the importation of used equipment associated with beekeeping

Competent Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the equipment:

   EITHER
   a) comes from a country or zone free from A. tumida infestation; and
   b) contains no live honey bees or bee brood;

   OR
   c) contains no live honey bees or bee brood; and
   d) has been thoroughly cleaned, and treated to ensure the destruction of A. tumida spp., in conformity with one of the procedures referred to in Chapter X.X. (under study);

   AND

2. all precautions have been taken to prevent infestation/contamination.

Article 9.4.9.

Recommendations for the importation of honey-bee collected pollen and beeswax (in the form of honeycomb)

Competent Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the products:

   EITHER
   a) comes from a country or zone free from A. tumida infestation; and
   b) contains no live honey bees or bee brood;

   OR
   c) contains no live honey bees or bee brood; and
   d) has been thoroughly cleaned, and treated to ensure the destruction of A. tumida spp., in conformity with one of the procedures referred to in Chapter X.X. (under study);
Annex XX (contd)

AND

2. all precautions have been taken to prevent infestation/contamination.

Article 9.4.10.

Recommendations for the importation of comb honey

Competent Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. comes from a country or zone free from A. tumida infestation; and
2. contains no live honey bees or bee brood;

OR

3. were subjected to a treatment at a temperature of -12°C or lower in the core of the product during at least 24 hours.

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CHAPTER 9.5.

TROPILAEELAPS INFESTATION OF HONEY BEES

Article 9.5.1.

General provisions

For the purposes of this Chapter, Tropilaelaps infestation of the honey bee Apis mellifera L. is caused by the mites Tropilaelaps dareae, T. komigereum, T. thaei and T. merodesae. The mite is an ectoparasite of brood of A. pis. mellifera L., A. pis laboriosa and A. pis dorsata, and cannot survive for periods of more than 7 days away from bee brood.

Early signs of infection normally go unnoticed, but the growth in the mite population is rapid leading to high hive mortality. The infection spreads by direct contact from adult bee to adult bee, and by the movement of infested bees and bee brood. The mite can also act as a vector for viruses of the honey bee.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 9.5.1.bis

Trade in commodities

When authorising import or transit of the following commodities, Veterinary Authorities should not require any Tropilaelaps infestation related conditions, regardless of the Tropilaelaps status of the honey bee population of the exporting country or zone:

1. honey bee semen, honey bee eggs and honey bee venom;
2. extracted honey and beeswax (not in the form of honeycomb).

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the Tropilaelaps status of the honey bee population of the exporting country or zone.

Article 9.5.2.

Determination of the Tropilaelaps status of a country or zone/ compartment

The Tropilaelaps status of a country or zone/ compartment (under study) can only be determined after considering the following criteria:

1. a risk assessment has been conducted, identifying all potential factors for Tropilaelaps occurrence and their historic perspective;
2. Tropilaelaps infestation should be notifiable in the whole country or zone/ compartment (under study) and all clinical signs suggestive of Tropilaelaps infestation should be subjected to field and laboratory investigations;
Annex XX (contd)

3. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of Tropilaelaps infestation;

4. the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees should have current knowledge of, and authority over, all domesticated apiaries in the country.

Article 9.5.3.

Country or zone/ compartment (under study) free from Tropilaelaps spp

1. Historically free status

A country or zone/compartment (under study) may be considered free from the disease after conducting a risk assessment as referred to in Article 9.5.2. but without formally applying a specific surveillance programme if the country or zone/compartment (under study) complies with the provisions of Chapter 1.4.

2. Free status as a result of an eradication programme

A country or zone/compartment (under study) which does not meet the conditions of point 1 above may be considered free from Tropilaelaps infestation after conducting a risk assessment as referred to in Article 9.5.2. and when:

a) the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees has current knowledge of, and authority over, all domesticated apiaries existing in the country or zone/compartment (under study);

b) Tropilaelaps infestation is notifiable in the whole country or zone/compartment (under study), and any clinical cases suggestive of Tropilaelaps infestation are subjected to field and laboratory investigations;

c) for the 3 years following the last reported case of Tropilaelaps infestation, an annual survey supervised by the Veterinary Authority, with negative results, have been carried out on a representative sample of apiaries in the country or zone/compartment (under study) to provide a confidence level of at least 95% of detecting Tropilaelaps infestation if at least 1% of the apiaries were infected at a within- apiary prevalence rate of at least 5% of the hives; such surveys may be targeted towards areas with a higher likelihood of infestation;

d) to maintain free status, an annual survey supervised by the Veterinary Authority, with negative results, is carried out on a representative sample of apiaries in the country or zone/compartment (under study) to indicate that there has been no new cases; such surveys may be targeted towards areas with a higher likelihood of disease;

e) (under study) there is no self-sustaining feral population of A. mellifera, A. dorsata or A. laboriosa, or other possible host species in the country or zone/compartment (under study);

f) the importation of the commodities listed in this Chapter into the country or zone/compartment (under study) is carried out, in conformity with the recommendations of this Chapter.
Article 9.5.4.

Recommendations on safe commodities

Regardless of the status of the exporting country with regard to Tropilaelaps infestation, Veterinary Authorities should authorise without restriction the import or transit through their territory of the following commodities:

1. honey bee semen, honey bee eggs and honey bee venom;
2. extracted honey and beeswax (not in the form of honeycomb).

Article 9.5.5.

Recommendations for the importation of live queen honey bees, worker bees and drones with associated brood combs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the bees come from a country or zone/compartment (under study) officially free from Tropilaelaps infestation.

Article 9.5.6.

Recommendations for the importation of live queen honey bees, worker bees and drones without associated brood combs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the bees have been held in isolation from brood and bees with access to brood, for a period of at least 7 days.

Article 9.5.7.

Recommendations for the importation of used equipment associated with beekeeping

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the equipment:

1. comes from a country or zone/compartment (under study) free from Tropilaelaps infestation; or
2. contains no live honey bees or bee brood and has been held away from contact with live honey bees for at least 7 days prior to shipment; or
3. has been treated to ensure the destruction of Tropilaelaps spp., in conformity with one of the procedures referred to in Chapter X.X. (under study).

Article 9.5.8.

Recommendations for the importation of honey-bee collected pollen, beeswax (in the form of honeycomb), comb honey and propolis

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:
Annex XX (contd)

1. come from a country or zone/compartment (under study) free from *Tropilaelaps* infestation; or

2. contain no live honey bees or bee brood and has been held away from contact with live honey bees for at least 7 days prior to shipment; or

3. have been treated to ensure the destruction of *Tropilaelaps* spp., in conformity with one of the procedures referred to in Chapter X.X. (under study).

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CHAPTER 9.6.

VARROOSIS OF HONEY BEES

Article 9.6.1.

General provisions

For the purposes of this Chapter, varroosis is a disease of the honey bee Apis mellifera L. It is caused by the Korea and Japan haplotypes of the mite Varroa destructor, the original hosts of which are the Korea and Japan haplotypes of Apis cerana (under study). The mite is an ectoparasite of adults and brood of Apis mellifera L. During its life cycle, sexual reproduction occurs inside the honey bee brood cells. Early signs of infection normally go unnoticed, and only when infection is heavy does it become apparent. The infection spreads by direct contact from adult bee to adult bee, and by the movement of infested bees and bee brood. The mite can also act as a vector for viruses of the honey bee.

The number of parasites steadily increases with increasing brood activity and the growth of the bee population, especially late in the season when clinical signs of infestation can first be recognised. The life span of an individual the mite depends on temperature and humidity but, in practice, it can be said to last from some days to a few months.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 9.6.1.bis

Trade in commodities

When authorising import or transit of the following commodities, Veterinary Authorities should not require any varroosis related conditions, regardless of the varroosis status of the honey bee population of the exporting country or zone:

1. honey bee semen, honey bee eggs and honey bee venom;
2. extracted honey and beeswax (not in the form of honey comb).

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the varroosis status of the honey bee population of the exporting country or zone.

Article 9.6.2.

Determination of the varroosis status of a country or zone/compartment

The varroosis status of a country or zone/compartment (under study) can only be determined after considering the following criteria:

1. a risk assessment has been conducted, identifying all potential factors for varroosis occurrence and their historic perspective;
2. varroosis should be notifiable in the whole country or zone/compartment (under study) and all clinical signs suggestive of varroosis should be subjected to field and laboratory investigations;
Annex XX (contd)

3. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of varroosis;

4. the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees should have current knowledge of, and authority over, all domesticated apiaries in the country.

Article 9.6.3.

Country or zone/compartment (under study) free from varroosis

1. Historically free status

A country or zone/compartment (under study) may be considered free from the disease after conducting a risk assessment as referred to in Article 9.6.2. but without formally applying a specific surveillance programme (historical freedom) if the country or zone/compartment (under study) complies with the provisions of Chapter 1.4.

2. Free status as a result of an eradication programme

A country or zone/compartment (under study) which does not meet the conditions of point 1 above may be considered free from varroosis after conducting a risk assessment as referred to in Article 9.6.2. and when:

a) the Veterinary Authority or other Competent Authority with responsibility for the health reporting and control of diseases of honey bees has current knowledge of, and authority over, all domesticated apiaries existing in the country or zone/compartment (under study);

b) varroosis is notifiable in the whole country or zone/compartment (under study), and any clinical cases suggestive of varroosis are subjected to field and laboratory investigations;

c) for the 3 years following the last reported case of varroosis, an annual survey supervised by the Veterinary Authority, with negative results, have been carried out on a representative sample of apiaries in the country or zone/compartment (under study) to provide a confidence level of at least 95% of detecting varroosis if at least 1% of the apiaries were infected at a within-apiary prevalence rate of at least 5% of the hives; such surveys may be targeted towards areas with a higher likelihood of disease;

d) to maintain free status, an annual survey supervised by the Veterinary Authority, with negative results, is carried out on a representative sample of apiaries in the country or zone/compartment (under study) to indicate that there has been no new cases; such surveys may be targeted towards areas with a higher likelihood of disease;

e) (under study) there is no self-sustaining feral population of A. mellifera, the Korea and Japan haplotypes of A. cerana or other possible host species in the country or zone/compartment (under study);

f) the importation of the commodities listed in this Chapter into the country or zone/compartment (under study) is carried out in conformity with the recommendations of this Chapter.
Recommendations on safe commodities

Regardless of the varroosis status of the exporting country, Veterinary Authorities should authorise without restriction the import or transit through their territory of the following commodities:

1. honey bee semen, honey bee eggs and honey bee venom;
2. extracted honey and beeswax (not in the form of honeycomb).

Recommendations for the importation of live queen honey bees, worker bees and drones with or without associated brood combs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the bees come from a country or zone/compartment (under study) officially free from varroosis.

Recommendations for the importation of larvae and pupae of honey bees

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. were sourced from a free country or zone/compartment (under study); or
2. have originated from queens in a quarantine station and were inspected and found free of Varroa destructor.

Recommendations for the importation of used equipment associated with beekeeping

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the equipment:

1. comes from a country or zone/compartment (under study) free from varroosis; or
2. contains no live honey bees or bee brood and has been held away from contact with live honey bees for at least 7 days prior to shipment; or
3. has been treated to ensure the destruction of Varroa destructor, in conformity with one of the procedures referred to in Chapter X.X. (under study).

Recommendations for the importation of honey-bee collected pollen, beeswax (in the form of honeycomb), comb honey and propolis
Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. come from a country or zone/compartment (under study) free from varroosis; or

2. contain no live honey bees or bee brood and has been held away from contact with live honey bees for at least 7 days prior to shipment; or

3. have been treated to ensure the destruction of Varroa destructor, in conformity with one of the procedures referred to in Chapter X.X. (under study).
CHAPTER 10.4.

AVIAN INFLUENZA

Article 10.4.1.

General provisions

1. For the purposes of international trade, avian influenza in its notifiable form (NAI) is defined as an infection of poultry caused by any influenza A virus of the H5 or H7 subtypes or by any AI virus with an intravenous pathogenicity index (IVPI) greater than 1.2 (or as an alternative at least 75% mortality) as described below. NAI viruses can be divided into highly pathogenic notifiable avian influenza (HPNAI) and low pathogenicity notifiable avian influenza (LPNAI):

a) HPNAI viruses have an IVPI in 6-week-old chickens greater than 1.2 or, as an alternative, cause at least 75% mortality in 4- to 8-week-old chickens infected intravenously. H5 and H7 viruses which do not have an IVPI of greater than 1.2 or cause less than 75% mortality in an intravenous lethality test should be sequenced to determine whether multiple basic amino acids are present at the cleavage site of the haemagglutinin molecule (HA0); if the amino acid motif is similar to that observed for other HPNAI isolates, the isolate being tested should be considered as HPNAI;

b) LPNAI are all influenza A viruses of H5 and H7 subtype that are not HPNAI viruses.

2. Poultry is defined as 'all domesticated birds, including backyard poultry, used for the production of meat or eggs for consumption, for the production of other commercial products, for restocking supplies of game, or for breeding these categories of birds, as well as fighting cocks used for any purpose'.

Birds that are kept in captivity for any reason other than those reasons referred to in the preceding paragraph, including those that are kept for shows, races, exhibitions, competitions or for breeding or selling these categories of birds as well as pet birds, are not considered to be poultry.

3. For the purposes of international trade, this Chapter deals not only with the occurrence of clinical signs caused by NAI virus, but also with the presence of infection with NAI virus in the absence of clinical signs.

4. For the purposes of international trade, a Member should not impose immediate trade bans on the trade in commercial poultry commodities in response to a notification, according to Article 1.2.3. of the Terrestrial Code, of infection with HPAI and LPAI virus in birds other than poultry, including wild birds, according to Article 1.2.3. of the Terrestrial Code.

5. Antibodies to H5 or H7 subtype of NAI virus, which have been detected in poultry and are not a consequence of vaccination, have to be further immediately investigated. In the case of isolated serological positive results, NAI infection may be ruled out on the basis of a thorough epidemiological and laboratory investigation that does not demonstrate further evidence of NAI infection.
6. The following defines the occurrence of infection with NAI virus:

   a) HPNAI virus has been isolated and identified as such or viral RNA specific for HPNAI has been detected in poultry or a product derived from poultry; or

   b) LPNAI virus has been isolated and identified as such or viral RNA specific for LPNAI has been detected in poultry or a product derived from poultry.

For the purposes of the Terrestrial Code, ‘NAI free establishment’ means an establishment in which the poultry have shown no evidence of NAI infection, based on surveillance in accordance with Articles 10.4.27. to 10.4.33.

For the purposes of the Terrestrial Code, the incubation period for NAI shall be 21 days.

Standards for diagnostic tests, including pathogenicity testing, are described in the Terrestrial Manual. Any vaccine used should comply with the standards described in the Terrestrial Manual.

Article 10.4.2.

**Determination of the NAI status of a country, zone or compartment**

The NAI status of a country, a zone or a compartment can be determined on the basis of the following criteria:

1. NAI is notifiable in the whole country, an on-going NAI awareness programme is in place, and all notified suspect occurrences of NAI are subjected to field and, where applicable, laboratory investigations;

2. appropriate surveillance is in place to demonstrate the presence of infection in the absence of clinical signs in poultry, and the risk posed by birds other than poultry; this may be achieved through a NAI surveillance programme in accordance with Articles 10.4.27. to 10.4.33.;

3. consideration of all epidemiological factors for NAI occurrence and their historical perspective.

Article 10.4.3.

**NAI free country, zone or compartment**

A country, zone or compartment may be considered free from NAI when it has been shown that neither HPNAI nor LPNAI infection in poultry has been present in the country, zone or compartment for the past 12 months, based on surveillance in accordance with Articles 10.4.27. to 10.4.33.

If infection has occurred in poultry in a previously free country, zone or compartment, NAI free status can be regained:

1. In the case of HPNAI infections, 3 months after a stamping-out policy (including disinfection of all affected establishments) is applied, providing that surveillance in accordance with Articles 10.4.27. to 10.4.33. has been carried out during that three-month period.
2. In the case of LPNAI infections, poultry may be kept for slaughter for human consumption subject to conditions specified in Articles 10.4.20. or 10.4.21. or a stamping-out policy may be applied; in either case, 3 months after the disinfection of all affected establishments, providing that surveillance in accordance with Articles 10.4.27. to 10.4.33. has been carried out during that three-month period.

Article 10.4.4.

**HPNAI free country, zone or compartment**

A country, zone or compartment may be considered free from HPNAI when:

1. it has been shown that HPNAI infection in poultry has not been present in the country, zone or compartment for the past 12 months, although its LPNAI status may be unknown; or

2. when, based on surveillance in accordance with Articles 10.4.27. to 10.4.33., it does not meet the criteria for freedom from NAI but any NAI virus detected has not been identified as HPNAI virus.

The surveillance may need to be adapted to parts of the country or existing zones or compartments depending on historical or geographical factors, industry structure, population data, or proximity to recent outbreaks.

If infection has occurred in poultry in a previously free country, zone or compartment, HPNAI free status can be regained 3 months after a stamping-out policy (including disinfection of all affected establishments) is applied, providing that surveillance in accordance with Articles 10.4.27. to 10.4.33. has been carried out during that three-month period.

Article 10.4.5.

**Recommendations for importation from a NAI free country, zone or compartment**

*for live poultry (other than day-old poultry)*

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the poultry showed no clinical sign of NAI on the day of shipment;

2. the poultry were kept in a NAI free country, zone or compartment since they were hatched or for at least the past 21 days;

3. the required surveillance, in accordance with Articles 10.4.27. to 10.4.33., has been carried out on the establishment within at least the past 21 days the poultry are transported in new or appropriately sanitized containers;

4. if vaccinated, the poultry have been vaccinated against NAI, it has been done in accordance with Articles 10.4.27. to 10.4.33., the provisions of the Terrestrial Manual and, in that case, the nature of the vaccine used and the date of vaccination should be have been attached to the certificate.
Annex XXI (contd)

Article 10.4.6.

Recommendations for the importation of live birds other than poultry

Regardles of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the birds showed no clinical sign of infection with a virus which would be considered NAI in poultry on the day of shipment;

2. the birds were kept in isolation approved by the Veterinary Services since they were hatched or for at least the 21 days prior to shipment and showed no clinical sign of infection with a virus which would be considered NAI in poultry during the isolation period;

3. a statistically valid sample of the birds at a design prevalence acceptable to the importing country was subjected to a diagnostic test within 14 days prior to shipment to demonstrate freedom from infection with a virus which would be considered NAI in poultry;

4. the birds are transported in new or appropriately sanitized containers;

5. if the birds have been vaccinated against NAI, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should be attached to the certificate.

Article 10.4.7.

Recommendations for importation from a NAI free country, zone or compartment for day-old live poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the poultry were kept in a NAI free country, zone or compartment since they were hatched;

2. the poultry were derived from parent flocks which had been kept in a NAI free country, zone or compartment for at least 21 days prior to and at the time of the collection of the eggs;

3. the poultry are transported in new or appropriately sanitized containers;

4. if the poultry or the parent flocks were vaccinated against NAI, vaccination was carried out in accordance with articles 10.4.27. to 10.4.33. the provisions of the Terrestrial Manual and, in that case, the nature of the vaccine used and the date of vaccination should be attached to the certificate.

Article 10.4.8.

Recommendations for importation from a HPNAI free country, zone or compartment for day-old live poultry
Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the poultry were kept in a HPNAI free country, zone or compartment since they were hatched;
2. the poultry were derived from parent flocks which had been kept in a NAI free establishment for at least 21 days prior to and at the time of the collection of the eggs;
3. the poultry are transported in new or appropriately sanitized containers;
4. if the poultry or the parent flocks were have been vaccinated against NAI, vaccination was carried out it has been done in accordance with articles 10.4.27. to 10.4.33. the provisions of the Terrestrial Manual and, in that case, the nature of the vaccine used and the date of vaccination should be have been attached to the certificate.

Article 10.4.9.

Recommendations for the importation of day-old live birds other than poultry

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the birds showed no clinical signs suggestive of NAI on the day of shipment;
2. the birds were hatched and kept in isolation approved by the Veterinary Services;
3. the parent flock birds were subjected to a diagnostic test at the time of the collection of the eggs to demonstrate freedom from infection with NAIV;
4. the birds are transported in new or appropriately sanitized containers;
5. if the birds or parent flocks were have been vaccinated against NAI, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should have been attached to the certificate.

Article 10.4.10.

Recommendations for importation from a NAI free country, zone or compartment

for hatching eggs of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the eggs came from a NAI free country, zone or compartment;
2. the eggs were derived from parent flocks which had been kept in a NAI free country, zone or compartment for at least 21 days prior to and at the time of the collection of the eggs;
3. the eggs are transported in new or appropriately sanitized containers.
Annex XXI (contd)

34. if the parent flocks were have been vaccinated against NAI, vaccination was carried out it has been done in accordance with Articles 10.4.27. to 10.4.33., the provisions of the Terrestrial Manual and, in that case, the nature of the vaccine used and the date of vaccination should be have been attached to the certificate.

4. the eggs are transported in new or appropriately sanitized containers.

Article 10.4.11.

Recommendations for importation from a HPNAI free country, zone or compartment

for hatching eggs of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the eggs came from a HPNAI free country, zone or compartment;

2. the eggs were derived from parent flocks which had been kept in a NAI free establishment for at least 21 days prior to and at the time of the collection of the eggs;

3. the eggs have had their surfaces sanitised (in accordance with Chapter 6.3.) and;

4. the eggs are transported in new or appropriately sanitized containers.

45. if the parent flocks were have been vaccinated against NAI, vaccination was carried out it has been done in accordance with Articles 10.4.27. to 10.4.33., the provisions of the Terrestrial Manual and, in that case, the nature of the vaccine used and the date of vaccination should be have been attached to the certificate.

Article 10.4.12.

Recommendations for the importation of hatching eggs from birds other than poultry

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the parent flock birds were subjected to a diagnostic test 7 days prior to and at the time of the collection of the eggs to demonstrate freedom from infection with NAIV;

2. the eggs have had their surfaces sanitized (in accordance with Chapter 6.3.) and;

3. the eggs are transported in new or appropriately sanitized containers.

4. the parent flocks have not been vaccinated against NAI; if the parent flocks were have been vaccinated against NAI, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should also be have been attached to the certificate.

Article 10.4.13.

Recommendations for importation from a NAI free country, zone or compartment
Recommendations for importation from a HPNAI free country, zone or compartment for eggs for human consumption

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:
1. the eggs were produced and packed in a HPNAI free country, zone or compartment;
2. the eggs have had their surfaces sanitized (in accordance with Chapter 6.3.) and;
3. the eggs are transported in new or appropriately sanitized packaging material containers.

Article 10.4.15.

Recommendations for importation from a NAI free country, zone or compartment for egg products

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the egg products come from, and were processed in, a NAI free country, zone or compartment.

Article 10.4.16.

Recommendations for importation of egg products of poultry from a country, zone or compartment not considered free from NAI

Regardless of the NAI status of the country of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:
1. the egg products are commodity derived from eggs which meet the requirements of Articles 10.4.12, 10.4.13, or 10.4.14.; or
2. the egg products were commodity has been processed to ensure the destruction of NAI virus in accordance with Article 10.4.25.;

AND

3. the necessary precautions were taken after processing to avoid contact of the commodity with any source of NAI virus;
4. the eggs are transported in new or appropriately sanitized packaging material.
Annex XXI (contd)

Article 10.4.17.

Recommendations for importation from a NAI free country, zone or compartment
for poultry semen

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the donor poultry:

1. showed no clinical sign of NAI on the day of semen collection;
2. were kept in a NAI free country, zone or compartment for at least the 21 days prior to and at the time of semen collection.

Article 10.4.18.

Recommendations for the importation from a HPNAI free country, zone or compartment
for poultry semen

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the donor poultry:

1. showed no clinical sign of HPNAI on the day of semen collection;
2. were kept in a HPNAI free country, zone or compartment for at least the 21 days prior to and at the time of semen collection.

Article 10.4.19.

Recommendations for the importation of semen of birds other than poultry

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the donor birds:

1. were kept in isolation approved by the Veterinary Services for at least the 21 days prior to semen collection;
2. showed no clinical sign of infection with a virus which would be considered NAI in poultry during the isolation period;
3. were tested within 14 days prior to semen collection and shown to be free of NAI infection.

Article 10.4.20.

Recommendations for importation from a NAI free country, zone or compartment
for fresh meat of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of fresh meat comes from poultry:
1. which have been kept in a NAI free country, zone or compartment since they were hatched or for at least the past 21 days;

2. which have been slaughtered in an approved abattoir in a NAI free country, zone or compartment and have been subjected to ante-mortem and post-mortem inspections in accordance with Chapter 6.2. and have been found free of any signs suggestive of NAI.

Article 10.4.21.

Recommendations for importation from a HPNAI free country, zone or compartment

for fresh meat of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of fresh meat comes from poultry:

1. which have been kept in a HPNAI free country, zone or compartment since they were hatched or for at least the past 21 days;

2. which have been slaughtered in an approved abattoir in a HPNAI free country, zone or compartment and have been subjected to ante-mortem and post-mortem inspections in accordance with Chapter 6.2. and have been found free of any signs suggestive of NAI.

Article 10.4.22.

Recommendations for the importation of meat products of poultry

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the commodity is derived from fresh meat which meet the requirements of Articles 10.4.20. or 10.4.21.; or

2. the commodity has been processed to ensure the destruction of avian influenza NAI virus in accordance with Article 10.4.26.;

AND

3. the necessary precautions were taken to avoid contact of the commodity with any source of NAI virus.

Article 10.4.23.

Recommendations for the importation of products of poultry origin intended for use in animal feeding, or for agricultural or industrial use other than feather meal

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities were processed in a NAI free country, zone or compartment from poultry which were kept in a NAI free country, zone or compartment from the time they were hatched until the time of slaughter or for at least the 21 days preceding slaughter; or
2. these commodities have been processed to ensure the destruction of avian influenza NAI virus (under study); AND
3. the necessary precautions were taken to avoid contact of the commodity with any source of NAI virus.

Article 10.4.24.

Recommendations for the importation of feathers and down of poultry

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities originated from poultry as described in Articles 10.4.20. or 10.4.21. and were processed in a NAI free country, zone or compartment from poultry which were kept in a NAI free country, zone or compartment from the time they were hatched until the time of slaughter or for at least the 21 days preceding slaughter; or
2. these commodities have been processed to ensure the destruction of avian influenza NAI virus (under study); AND
3. the necessary precautions were taken to avoid contact of the commodity with any source of avian influenza NAI virus.

Article 10.4.24 bis.

Recommendations for the importation of feathers and down of birds other than poultry

Regardless of the NAI status of the country of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities have been processed to ensure the destruction of NAI virus (under study); and
2. the necessary precautions were taken to avoid contact of the commodity with any source of NAI virus.

Article 10.4.24 bis.

Recommendations for the importation of feather meal

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities were processed in a NAI free country, zone or compartment from poultry which were kept in a NAI free country, zone or compartment from the time they were hatched until the time of slaughter or for at least the 21 days preceding slaughter; or
2. these commodities have been processed either:
a) **moist heat** at a minimum temperature of 118°C for a minimum of 40 minutes; or

b) with a continuous hydrolysing process under at least 3.79 bar of pressure with steam at a minimum temperature of 122°C for a minimum of 15 minutes;

3. the necessary precautions were taken to avoid contact of the commodity with any source of **avian influenza NAI virus**.

**Article 10.4.25.**

**Procedures for the inactivation of the AI virus in eggs and egg products**

The following times for industry standard temperatures are suitable for the inactivation of HPNAI virus present in eggs and egg products:

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whole egg</td>
<td>60</td>
</tr>
<tr>
<td>Whole egg blends</td>
<td>60</td>
</tr>
<tr>
<td>Whole egg blends</td>
<td>61.1</td>
</tr>
<tr>
<td>Liquid egg white</td>
<td>55.6</td>
</tr>
<tr>
<td>Liquid egg white</td>
<td>56.7</td>
</tr>
<tr>
<td>10% salted yolk</td>
<td>62.2</td>
</tr>
<tr>
<td>Dried egg white</td>
<td>67</td>
</tr>
<tr>
<td>Dried egg white</td>
<td>54.4</td>
</tr>
</tbody>
</table>

The listed temperatures are indicative of a range that achieves a 7-log kill. Where scientifically documented, variances from these times and temperatures may also be suitable when they achieve the inactivation of the virus.

**Article 10.4.26.**

**Procedures for the inactivation of the AI virus in meat**

A procedure which produces a core temperature of 70°C for 3.5 seconds is suitable for the inactivation of HPNAI virus present in meat.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poultry meat</td>
<td>60.0</td>
</tr>
<tr>
<td></td>
<td>65.0</td>
</tr>
<tr>
<td></td>
<td>70.0</td>
</tr>
<tr>
<td></td>
<td>73.9</td>
</tr>
</tbody>
</table>
Annex XXI (contd)

Article 10.4.27.

Surveillance: introduction

Articles 10.4.27. to 10.4.33. define the principles and provide a guide on the surveillance of NAI complementary to Chapter 1.4., applicable to Members seeking to determine their NAI status. This may be for the entire country, zone or compartment. Guidance for Members seeking free status following an outbreak and for the maintenance of NAI status is also provided.

The presence of avian influenza viruses in wild birds creates a particular problem. In essence, no Member can declare itself free from avian influenza (AI) in wild birds. However, the definition of NAI in this Chapter refers to the infection in poultry only, and Articles 10.4.27. to 10.4.33. were developed under this definition.

The impact and epidemiology of NAI differ widely in different regions of the world and therefore it is impossible to provide specific recommendations for all situations. It is axiomatic that the surveillance strategies employed for demonstrating freedom from NAI at an acceptable level of confidence will need to be adapted to the local situation. Variables such as the frequency of contacts of poultry with wild birds, different biosecurity levels and production systems and the commingling of different susceptible species including domestic waterfowl require specific surveillance strategies to address each specific situation. It is incumbent upon the Member to provide scientific data that explains the epidemiology of NAI in the region concerned and also demonstrates how all the risk factors are managed. There is therefore considerable latitude available to Members to provide a well-reasoned argument to prove that absence of NAI virus (NAIV) infection is assured at an acceptable level of confidence.

Surveillance for NAI should be in the form of a continuing programme designed to establish that the country, zone or compartment, for which application is made, is free from NAIV infection.

Article 10.4.28.

Surveillance: general conditions and methods

1. A surveillance system in accordance with Chapter 1.4. should be under the responsibility of the Veterinary Authority. In particular:

   a) a formal and ongoing system for detecting and investigating outbreaks of disease or NAI infection should be in place;

   b) a procedure should be in place for the rapid collection and transport of samples from suspect cases of NAI to a laboratory for NAI diagnosis as described in the Terrestrial Manual;

   c) a system for recording, managing and analysing diagnostic and surveillance data should be in place.

2. The NAI surveillance programme should:
a) include an early warning system throughout the production, marketing and processing chain for reporting suspicious cases. Farmers and workers, who have day-to-day contact with poultry, as well as diagnosticians, should report promptly any suspicion of NAI to the Veterinary Authority. They should be supported directly or indirectly (e.g. through private veterinarians or veterinary para-professionals) by government information programmes and the Veterinary Authority. All suspected cases of NAI should be investigated immediately. As suspicion cannot always be resolved by epidemiological and clinical investigation alone, samples should be taken and submitted to a laboratory for appropriate tests. This requires that sampling kits and other equipment are available for those responsible for surveillance. Personnel responsible for surveillance should be able to call for assistance from a team with expertise in NAI diagnosis and control. In cases where potential public health implications are suspected, notification to the appropriate public health authorities is essential;

b) implement, when relevant, regular and frequent clinical inspection, serological and virological testing of high-risk groups of animals, such as those adjacent to a NAI infected country, zone or compartment, places where birds and poultry of different origins are mixed, such as live bird markets, poultry in close proximity to waterfowl or other potential sources of NAIV.

An effective surveillance system will periodically identify suspicious cases that require follow-up and investigation to confirm or exclude that the cause of the condition is NAIV. The rate at which such suspicious cases are likely to occur will differ between epidemiological situations and cannot therefore be predicted reliably. Applications for freedom from NAIV infection should, in consequence, provide details of the occurrence of suspicious cases and how they were investigated and dealt with. This should include the results of laboratory testing and the control measures to which the animals concerned were subjected during the investigation (quarantine, movement stand-still orders, etc.).

Article 10.4.29.

Surveillance strategies

1. Introduction

The target population for surveillance aimed at identification of disease and infection should cover all the susceptible poultry species within the country, zone or compartment. Active and passive surveillance for NAIV should be ongoing. The frequency of active surveillance should be at least every 6 months. Surveillance should be composed of random and targeted approaches using virological, serological and clinical methods.

The strategy employed may be based on randomised sampling requiring surveillance consistent with demonstrating the absence of NAIV infection at an acceptable level of confidence. The frequency of sampling should be dependent on the epidemiological situation. Random surveillance is conducted using serological tests described in the Terrestrial Manual. Positive serological results should be followed up with virological methods.

Targeted surveillance (e.g. based on the increased likelihood of infection in particular localities or species) may be an appropriate strategy. Virological and serological methods should be used concurrently to define the NAI status of high risk populations.
A Member should justify the surveillance strategy chosen as adequate to detect the presence of NAIV infection in accordance with Chapter 1.4 and the prevailing epidemiological situation, including cases of HPNAI detected in any birds. It may, for example, be appropriate to target clinical surveillance at particular species likely to exhibit clear clinical signs (e.g. chickens). Similarly, virological and serological testing could be targeted to species that may not show clinical signs (e.g. ducks).

If a Member wishes to declare freedom from NAIV infection in a specific zone or compartment, the design of the survey and the basis for the sampling process would need to be aimed at the population within the zone or compartment.

For random surveys, the design of the sampling strategy will need to incorporate epidemiologically appropriate design prevalence. The sample size selected for testing will need to be large enough to detect infection if it were to occur at a predetermined minimum rate. The sample size and expected disease prevalence determine the level of confidence in the results of the survey. The Member must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Chapter 1.4. Selection of the design prevalence in particular clearly needs to be based on the prevailing or historical epidemiological situation.

Irrespective of the survey approach selected, the sensitivity and specificity of the diagnostic tests employed are key factors in the design, sample size determination and interpretation of the results obtained. Ideally, the sensitivity and specificity of the tests used should be validated for the vaccination/infection history and the different species in the target population.

Irrespective of the testing system employed, surveillance system design should anticipate the occurrence of false positive reactions. If the characteristics of the testing system are known, the rate at which these false positives are likely to occur can be calculated in advance. There needs to be an effective procedure for following up positives to ultimately determine with a high level of confidence, whether they are indicative of infection or not. This should involve both supplementary tests and follow-up investigation to collect diagnostic material from the original sampling unit as well as flocks which may be epidemiologically linked to it.

The principles involved in surveillance for disease/infection are technically well defined. The design of surveillance programmes to prove the absence of NAIV infection/circulation needs to be carefully followed to avoid producing results that are either insufficiently reliable, or excessively costly and logistically complicated. The design of any surveillance programme, therefore, requires inputs from professionals competent and experienced in this field.

2. Clinical surveillance

Clinical surveillance aims at the detection of clinical signs of NAI at the flock level. Whereas significant emphasis is placed on the diagnostic value of mass serological screening, surveillance based on clinical inspection should not be underrated. Monitoring of production parameters, such as increased mortality, reduced feed and water consumption, presence of clinical signs of a respiratory disease or a drop in egg production, is important for the early detection of NAIV infection. In some cases, the only indication of LPNAIV infection may be a drop in feed consumption or egg production.

Clinical surveillance and laboratory testing should always be applied in series to clarify the status of NAI suspects detected by either of these complementary diagnostic approaches. Laboratory testing may confirm clinical suspicion, while clinical surveillance may contribute to confirmation of positive serology. Any sampling unit within which suspicious animals are detected should be classified as infected, have restrictions imposed upon it until evidence to the contrary is produced. NAIV infection is ruled out.
Identification of suspect flocks is vital to the identification of sources of NAIV and to enable the molecular, antigenic and other biological characteristics of the virus to be determined. It is essential that NAIV isolates are sent regularly to the regional Reference Laboratory for genetic and antigenic characterization.

3. **Virological surveillance**

Virological surveillance using tests described in the Terrestrial Manual should be conducted:

a) to monitor at risk populations;

b) to confirm clinically suspect cases;

c) to follow up positive serological results;

d) to test ‘normal’ daily mortality, to ensure early detection of infection in the face of vaccination or in establishments epidemiologically linked to an outbreak.

4. **Serological surveillance**

Serological surveillance aims at the detection of antibodies against NAIV. Positive NAIV antibody test results can have four possible causes:

a) natural infection with NAIV;

b) vaccination against NAIV;

c) maternal antibodies derived from a vaccinated or infected parent flock are usually found in the yolk and can persist in progeny for up to 4 weeks;

d) false positive results due to the lack of specificity of the test.

It may be possible to use serum collected for other survey purposes for NAI surveillance. However, the principles of survey design described in these recommendations and the requirement for a statistically valid survey for the presence of NAIV should not be compromised.

The discovery of clusters of seropositive flocks may reflect any of a series of events, including but not limited to the demographics of the population sampled, vaccinal exposure or infection. As clustering may signal infection, the investigation of all instances must be incorporated in the survey design. Clustering of positive flocks is always epidemiologically significant and therefore should be investigated.

If vaccination cannot be excluded as the cause of positive serological reactions, diagnostic methods to differentiate antibodies due to infection or vaccination should be employed.

The results of random or targeted serological surveys are important in providing reliable evidence that no NAIV infection is present in a country, zone or compartment. It is therefore essential that the survey be thoroughly documented.
Annex XXI (contd)

5. Virological and serological surveillance in vaccinated populations

The surveillance strategy is dependent on the type of vaccine used. The protection against AI is haemagglutinin subtype specific. Therefore, two broad vaccination strategies exist: 1) inactivated whole AI viruses, and 2) haemagglutinin expression-based vaccines.

In the case of vaccinated populations, the surveillance strategy should be based on virological and/or serological methods and clinical surveillance. It may be appropriate to use sentinel birds for this purpose. These birds should be unvaccinated, AI virus antibody free birds and clearly and permanently identified. Sentinel birds should be used only if no appropriate laboratory procedures are available. The interpretation of serological results in the presence of vaccination is described in Article 10.4.33.

Article 10.4.30.

Documentation of NAI or HPNAI free status

1. Members declaring freedom from NAI or HPNAI for the country, zone or compartment: additional surveillance procedures

In addition to the general conditions described in above mentioned articles, a Member declaring freedom from NAI or HPNAI for the entire country, or a zone or a compartment should provide evidence for the existence of an effective surveillance programme. The strategy and design of the surveillance programme will depend on the prevailing epidemiological circumstances and should be planned and implemented according to general conditions and methods described in this Chapter, to demonstrate absence of NAIV or HPNAIV infection, during the preceding 12 months in susceptible poultry populations (vaccinated and non-vaccinated). This requires the support of a laboratory able to undertake identification of NAIV or HPNAIV infection through virus detection and antibody tests described in the Terrestrial Manual. This surveillance may be targeted to poultry population at specific risks linked to the types of production, possible direct or indirect contact with wild birds, multi-age flocks, local trade patterns including live bird markets, use of possibly contaminated surface water, and the presence of more than one species on the holding and poor biosecurity measures in place.

2. Additional requirements for countries, zones or compartments that practise vaccination

Vaccination to prevent the transmission of HPNAI virus may be part of a disease control programme. The level of flock immunity required to prevent transmission will depend on the flock size, composition (e.g. species) and density of the susceptible poultry population. It is therefore impossible to be prescriptive. The vaccine must also comply with the provisions stipulated for NAI vaccines in the Terrestrial Manual. Based on the epidemiology of NAI in the country, zone or compartment, it may be that a decision is reached to vaccinate only certain species or other poultry subpopulations.

In all vaccinated flocks there is a need to perform virological and serological tests to ensure the absence of virus circulation. The use of sentinel poultry may provide further confidence of the absence of virus circulation. The tests have to be repeated at least every 6 months or at shorter intervals according to the risk in the country, zone or compartment.

Evidence to show the effectiveness of the vaccination programme should also be provided.
Article 10.4.31.

Countries, zones or compartments declaring that they have regained freedom from NAI or HPNAI following an outbreak: additional surveillance procedures

In addition to the general conditions described in the above-mentioned articles, a Member declaring that it has regained country, zone or compartment freedom from NAI or HPNAI virus infection should show evidence of an active surveillance programme depending on the epidemiological circumstances of the outbreak to demonstrate the absence of the infection. This will require surveillance incorporating virus detection and antibody tests described in the Terrestrial Manual. The use of sentinel birds may facilitate the interpretation of surveillance results.

A Member declaring freedom of country, zone or compartment after an outbreak of NAI or HPNAI (with or without vaccination) should report the results of an active surveillance programme in which the NAI or HPNAI susceptible poultry population undergoes regular clinical examination and active surveillance planned and implemented according to the general conditions and methods described in these recommendations. The surveillance should at least give the confidence that can be given by a randomized representative sample of the populations at risk.

Article 10.4.32.

NAI free establishments within HPNAI free compartments: additional surveillance procedures

The declaration of NAI free establishments requires the demonstration of absence of NAIV infection. Birds in these establishments should be randomly tested using virus detection or isolation tests, and serological methods, following the general conditions of these recommendations. The frequency of testing should be based on the risk of infection and at a maximum interval of 21 days.

Article 10.4.33.

The use and interpretation of serological and virus detection tests

Poultry infected with NAI virus produce antibodies to haemagglutinin (HA), neuraminidase (NA), nonstructural proteins (NSPs), nucleoprotein/matrix (NP/M) and the polymerase complex proteins. Detection of antibodies against the polymerase complex proteins will not be covered in this Chapter.

Tests for NP/M antibodies include direct and blocking ELISA, and agar gel immunodiffusion (AGID) tests. Tests for antibodies against NA include the neuraminidase inhibition (NI), indirect fluorescent antibody and direct and blocking ELISA tests. For the HA, antibodies are detected in haemagglutination inhibition (HI), ELISA and neutralization (SN) tests. The HI test is reliable in avian species but not in mammals. The SN test can be used to detect subtype specific antibodies to the haemagglutinin and is the preferred test for mammals and some avian species. The AGID test is reliable for detection of NP/M antibodies in chickens and turkeys, but not in other avian species. As an alternative, blocking ELISA tests have been developed to detect NP/M antibodies in all avian species.

The HI and NI tests can be used to subtype AI viruses into 16 haemagglutinin and 9 neuraminidase subtypes. Such information is helpful for epidemiological investigations and in categorization of AI viruses.

Poultry can be vaccinated with a variety of AI vaccines including inactivated whole AI virus vaccines, and haemagglutinin expression-based vaccines. Antibodies to the haemagglutinin confer subtype specific protection. Various strategies can be used to differentiate vaccinated from infected birds including serosurveillance in unvaccinated sentinel birds or specific serological tests in the vaccinated birds.
Annex XXI (contd)

AI virus infection of unvaccinated birds including sentinels is detected by antibodies to the NP/M, subtype specific HA or NA proteins, or NSP. Poultry vaccinated with inactivated whole AI vaccines containing an influenza virus of the same H sub-type but with a different neuraminidase may be tested for field exposure by applying serological tests directed to the detection of antibodies to the NA of the field virus. For example, birds vaccinated with H7N1 in the face of a H7N1 epidemic may be differentiated from infected birds (DIVA) by detection of subtype specific NA antibodies of the N1 protein of the field virus. Alternatively, in the absence of DIVA, inactivated vaccines may induce low titres of antibodies to NSP and the titre in infected birds would be markedly higher. Encouraging results have been obtained experimentally with this system, but it has not yet been validated in the field. In poultry vaccinated with haemagglutinin expression-based vaccines, antibodies are detected to the specific HA, but not any of the other AI viral proteins. Infection is evident by antibodies to the NP/M or NSP, or the specific NA protein of the field virus. Vaccines used should comply with the standards of the Terrestrial Manual.

All flocks with seropositive results should be investigated. Epidemiological and supplementary laboratory investigation results should document the status of NA1 infection/circulation for each positive flock.

A confirmatory test should have a higher specificity than the screening test and sensitivity at least equivalent than that of the screening test.

Information should be provided on the performance characteristics and validation of tests used.

1. The follow-up procedure in case of positive test results if vaccination is used

   In case of vaccinated populations, one has to exclude the likelihood that positive test results are indicative of virus circulation. To this end, the following procedure should be followed in the investigation of positive serological test results derived from surveillance conducted on NAI-vaccinated poultry. The investigation should examine all evidence that might confirm or refute the hypothesis that the positive results to the serological tests employed in the initial survey were not due to virus circulation. All the epidemiological information should be substantiated, and the results should be collated in the final report.

   Knowledge of the type of vaccine used is crucial in developing a serological based strategy to differentiate infected from vaccinated animals.

   a) Inactivated whole AI virus vaccines can use either homologous or heterologous neuraminidase subtypes between the vaccine and field strains. If poultry in the population have antibodies to NP/M and were vaccinated with inactivated whole AI virus vaccine, the following strategies should be applied:

      i) sentinel birds should remain NP/M antibody negative. If positive for NP/M antibodies, indicating AI virus infection, specific HI tests should be performed to identify H5 or H7 AI virus infection;

      ii) if vaccinated with inactivated whole AI virus vaccine containing homologous NA to field virus, the presence of antibodies to NSP could be indicative of infection. Sampling should be initiated to exclude the presence of NAIV by either virus isolation or detection of virus specific genomic material or proteins;

      iii) if vaccinated with inactivated whole AI virus vaccine containing heterologous NA to field virus, presence of antibodies to the field virus NA or NSP would be indicative of infection. Sampling should be initiated to exclude the presence of NAIV by either virus isolation or detection of virus specific genomic material or proteins.
b) Haemagglutinin expression-based vaccines contain the HA protein or gene homologous to the HA of the field virus. Sentinel birds as described above can be used to detect AI infection. In vaccinated or sentinel birds, the presence of antibodies against NP/M, NSP or field virus NA is indicative of infection. Sampling should be initiated to exclude the presence of NAIV by either virus isolation or detection of virus specific genomic material or proteins.

2. The follow-up procedure in case of positive test results indicative of infection for determination of infection due to HPNAI or LPNAI virus

The detection of antibodies indicative of a NAI virus infection as indicated in point a)i) above will result in the initiation of epidemiological and virological investigations to determine if the infections are due to HPNAI or LPNAI viruses.

Virological testing should be initiated in all antibody-positive and at risk populations. The samples should be evaluated for the presence of AI virus, by virus isolation and identification, and/or detection of influenza A specific proteins or nucleic acids (Figure 2). Virus isolation is the gold standard for detecting infection by AI virus and the method is described in the Terrestrial Manual. All AI virus isolates should be tested to determine HA and NA subtypes, and in vivo tested in chickens and/or sequencing of HA proteolytic cleavage site of H5 and H7 subtypes for determination of classification as HPNAI, LPNAI or LPAI (not notifiable) viruses. As an alternative, nucleic acid detection tests have been developed and validated; these tests have the sensitivity of virus isolation, but with the advantage of providing results within a few hours. Samples with detection of H5 and H7 HA subtypes by nucleic acid detection methods should either be submitted for virus isolation, identification, and in vivo testing in chickens, or sequencing of nucleic acids for determination of proteolytic cleavage site as HPNAI or LPNAI viruses. The antigen detection systems, because of low sensitivity, are best suited for screening clinical field cases for infection by Type A influenza virus looking for NP/M proteins. NP/M positive samples should be submitted for virus isolation, identification and pathogenicity determination.

Laboratory results should be examined in the context of the epidemiological situation. Corollary information needed to complement the serological survey and assess the possibility of viral circulation includes but is not limited to:

a) characterization of the existing production systems;

b) results of clinical surveillance of the suspects and their cohorts;

c) quantification of vaccinations performed on the affected sites;

d) sanitary protocol and history of the affected establishments;

e) control of animal identification and movements;

f) other parameters of regional significance in historic NAIV transmission.

The entire investigative process should be documented as standard operating procedure within the epidemiological surveillance programme.
Fig. 1. Schematic representation of laboratory tests for determining evidence of NAI infection through or following serological surveys.
Fig. 2. Schematic representation of laboratory tests for determining evidence of NAI infection using virological methods

The above diagram indicates the tests which are recommended for use in the investigation of poultry flocks.

<table>
<thead>
<tr>
<th>Key</th>
<th>Description</th>
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<tbody>
<tr>
<td>AGID</td>
<td>Agar gel immunodiffusion</td>
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<tr>
<td>DIVA</td>
<td>Differentiating infected from vaccinated animals</td>
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<tr>
<td>ELISA</td>
<td>Enzyme-linked immunosorbant assay</td>
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<tr>
<td>HA</td>
<td>Haemagglutinin</td>
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<tr>
<td>HI</td>
<td>Haemagglutination inhibition</td>
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<td>NA</td>
<td>Neuraminidase</td>
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<tr>
<td>NP/M</td>
<td>Nucleoprotein and matrix protein</td>
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<tr>
<td>NSP</td>
<td>Nonstructural protein</td>
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<td>S</td>
<td>No evidence of NAIV</td>
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CHAPTER 10.13.

NEWCASTLE DISEASE

Article 10.13.1.

General provisions

1. For the purposes of international trade, Newcastle disease (ND) is defined as an infection of poultry caused by a virus (NDV) of avian paramyxovirus serotype 1 (APMV-1) that meets one of the following criteria for virulence:

   a) the virus has an intracerebral pathogenicity index (ICPI) in day-old chicks (Gallus gallus) of 0.7 or greater; or

   b) multiple basic amino acids have been demonstrated in the virus (either directly or by deduction) at the C-terminus of the F2 protein and phenylalanine at residue 117, which is the N-terminus of the F1 protein. The term ‘multiple basic amino acids’ refers to at least three arginine or lysine residues between residues 113 and 116. Failure to demonstrate the characteristic pattern of amino acid residues as described above would require characterisation of the isolated virus by an ICPI test.

In this definition, amino acid residues are numbered from the N-terminus of the amino acid sequence deduced from the nucleotide sequence of the F0 gene, 113–116 corresponds to residues –4 to –1 from the cleavage site.

2. Poultry is defined as ‘all domesticated birds, including backyard poultry, used for the production of meat or eggs for consumption, for the production of other commercial products, for restocking supplies of game, or for breeding these categories of birds, as well as fighting cocks used for any purpose’.

Birds that are kept in captivity for any reason other than those reasons referred to in the preceding paragraph, including those that are kept for shows, races, exhibitions, competitions, or for breeding or selling these categories of birds as well as pet birds, are not considered to be poultry.

3. This Chapter deals with NDV infection of poultry as defined in point 2 above, in the presence or absence of clinical signs. For the purposes of international trade, a Member should not impose immediate trade bans on the trade in commercial poultry commodities in response to reports of a notification Article 1.2.3. of the Terrestrial Code of infection with NDV in birds other than poultry, including wild birds, according to Article 1.2.3. of the Terrestrial Code.

4. The occurrence of infection with NDV is defined as the isolation and identification of NDV as such or the detection of viral RNA specific for NDV.

5. For the purposes of the Terrestrial Code, the incubation period for ND shall be 21 days.

Standards for diagnostic tests, including pathogenicity testing, are described in the Terrestrial Manual. When the use of ND vaccines is appropriate, those vaccines should comply with the standards described in the Terrestrial Manual.
Article 10.13.2.

Determination of the ND status of a country, zone or compartment

The ND status of a country, a zone or a compartment can be determined on the basis of the following criteria:

1. ND is notifiable in the whole country, an on-going ND awareness programme is in place, and all notified suspect occurrences of ND are subjected to field and, where applicable, laboratory investigations;

2. appropriate surveillance is in place to demonstrate the presence of NDV infection in the absence of clinical signs in poultry, this may be achieved through an ND surveillance programme in accordance with Articles 10.13.20. to 10.13.24.;

3. consideration of all epidemiological factors for ND occurrence and their historical perspective.

Article 10.13.3.

ND free country, zone or compartment

A country, zone or compartment may be considered free from ND when it has been shown that NDV infection in poultry has not been present in the country, zone or compartment for the past 12 months, based on surveillance in accordance with Articles 10.13.20. to 10.13.24.

If infection has occurred in poultry in a previously free country, zone or compartment, ND free status can be regained three months after a stamping-out policy (including disinfection of all affected establishments) is applied, providing that surveillance in accordance with Articles 10.13.20. to 10.13.24. has been carried out during that three-month period.

Article 10.13.4.

Recommendations for importation from an ND free country, zone or compartment as defined in Article 10.13.3.

for live poultry (other than day-old poultry)

Veterinary authorities should require the presentation of an international veterinary certificate attesting that:

1. the poultry showed no clinical sign suggestive of ND on the day of shipment;

2. the poultry were kept in an ND free country, zone or compartment since they were hatched or for at least the past 21 days;

3. the poultry are transported in new or appropriately sanitized containers;

4. if the birds poultry have been vaccinated against ND, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should be have been attached to the certificate.
Recommendations for the importation of live birds other than poultry

Regardless of the ND status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the birds showed no clinical sign suggestive of ND on the day of shipment;
2. the birds were kept in isolation approved by the Veterinary Services since they were hatched or for at least the 21 days prior to shipment and showed no clinical sign of infection during the isolation period;
3. a statistically valid sample of the birds at a design prevalence acceptable to the importing country were subjected to a diagnostic test within 14 days prior to shipment to demonstrate freedom from infection with NDV;
4. the birds are transported in new or appropriately sanitized containers;
5. if the birds were vaccinated against ND, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should also be attached to the certificate.

Recommendations for importation from an ND free country, zone or compartment as defined in Article 10.13.3, for day-old live poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the poultry were hatched and kept in an ND free country, zone or compartment since they were hatched;
2. the poultry were derived from parent flocks which had been kept in an ND free country, zone or compartment for at least 21 days prior to and at the time of the collection of the eggs;
3. the poultry are transported in new or appropriately sanitized containers;
4. if poultry or parent flocks were vaccinated against ND, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should also be attached to the certificate.

Recommendations for the importation of day-old live birds other than poultry

Regardless of the ND status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the birds showed no clinical sign suggestive of ND on the day of shipment;
2. the birds were hatched and kept in isolation approved by the Veterinary Services;
Annex XXII (contd)

3. the parent flock birds were subjected to a diagnostic test at the time of the collection of the eggs to demonstrate freedom from infection with NDV;

4. the birds are transported in new or appropriately sanitized containers.

5. If the birds or parent flocks were have been vaccinated against ND, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should also be have been attached to the certificate.

Article 10.13.8.

Recommendations for importation from an ND free country, zone or compartment as defined in Article 10.13.3.

for hatching eggs of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the birds:

1. the eggs came from an ND free country, zone or compartment;

2. the eggs were derived from parent flocks which had been kept in an ND free country, zone or compartment for at least 21 days prior to and at the time of the collection of the eggs;

3. the eggs are transported in new or appropriately sanitized packing material containers.

4. If parent flocks were have been vaccinated against ND, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should also be have been attached to the certificate.

Article 10.13.9.

Recommendations for the importation of hatching eggs from birds other than poultry

Regardless of the ND status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the parent flock birds were subjected to a diagnostic test 7 days prior to and at the time of the collection of the eggs to demonstrate freedom from infection with NDV;

2. the eggs have had their surfaces sanitized (in accordance with Chapter 6.3.) and ;

3. the eggs are transported in new or appropriately sanitized packing material containers.

4. If the parent flocks were have been vaccinated against ND, it has been done in accordance with the provisions of the Terrestrial Manual and the nature of the vaccine used and the date of vaccination should also be have been attached to the certificate.
Article 10.13.10.

Recommendations for importation from an ND free country, zone or compartment as defined in Article 10.13.3. for eggs for human consumption

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the eggs were produced and packed in an ND free country, zone or compartment;
2. the eggs are transported in new or appropriately sanitized packing material containers.

Article 10.13.11.

Recommendations for importation from an ND free country, zone or compartment as defined in Article 10.13.3. for egg products

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the egg products come from, and were processed in, an ND free country, zone or compartment;
2. the egg products are transported in new or appropriately sanitized containers.

Article 10.13.12.

Recommendations for importation of egg products of poultry from a country, zone or compartment not considered free from ND for egg products

Regardless of the ND status of the country of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the commodity is derived from eggs which meet the requirements of Article 10.13.10.; or
2. the egg products are commodity has been processed to ensure the destruction of NDV (under study) in accordance with Article 10.13.19. and/or (under study);

AND

23. the necessary precautions were taken after processing to avoid contact of the egg products with any source of NDV;
3. the egg products are transported in new or appropriately sanitized containers.
Article 10.13.13.

Recommendations for importation from an ND free country, zone or compartment as defined in Article 10.13.3. for poultry semen

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the donor poultry:

1. showed no clinical sign suggestive of ND on the day of semen collection;
2. were kept in an ND free country, zone or compartment for at least the 21 days prior to and at the time of semen collection.


Recommendations for the importation of semen of birds other than poultry

Regardless of the ND status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the donor birds:

1. were kept in isolation approved by the Veterinary Services for at least the 21 days prior to and on the day of semen collection;
2. showed no clinical sign suggestive of infection with NDV during the isolation period and on the day of semen collection;
3. were subjected to a diagnostic test within 14 days prior to semen collection to demonstrate freedom from infection with NDV.

Article 10.13.15.

Recommendations for importation from an ND free country, zone or compartment as defined in Article 10.13.3. for fresh meat of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of fresh meat comes from poultry:

1. which have been kept in an ND free country, zone or compartment since they were hatched or for at least the past 21 days;
2. which have been slaughtered in an approved abattoir in an ND free country, zone or compartment and have been subjected to ante-mortem and post-mortem inspections in accordance with Chapter 6.2. and have been found free of any sign suggestive of ND.

Article 10.13.16.

Recommendations for importation from an ND free country, zone or compartment
Annex XXII (contd)

for meat products of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the commodity is derived from fresh meat which meets the requirements of Article 10.13.15. and has been processed in an ND free country, zone or compartment;

2. the necessary precautions were taken to avoid contact of the commodity with any source of NDV.

Article 10.13.17.

Recommendations for importation of meat products of poultry from a country, zone or compartment not considered free from ND

for meat products of poultry

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the entire consignment of meat comes from poultry which have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections in accordance with Chapter 6.2. and have been found free of any signs suggestive of ND; the commodity is derived from fresh meat which meet the requirements of Article 10.13.15.; or

2. the commodity has been processed to ensure the destruction of NDV (under study) in accordance with Article 10.13.19 quads quints (under study);

AND

3. the necessary precautions were taken to avoid contact of the commodity with any source of NDV.

Article 10.13.18.

Recommendations for the importation of products of poultry origin intended for use in animal feeding, or for agricultural or industrial use other than feather meal

Regardless of the ND status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities were processed in a ND free country, zone or compartment from poultry which were kept in a ND free country, zone or compartment from the time they were hatched until the time of slaughter or at least the 21 days preceding slaughter; or

2. these commodities have been processed to ensure the destruction of NDV (under study);

AND

3. the necessary precautions were taken to avoid contact of the commodity with any source of NDV.

Article 10.13.19.

Recommendations for the importation of feathers and down of poultry
Regardless of the ND status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities originated from poultry as described in Article 10.13.15. and were processed in a ND free country, zone or compartment from poultry which were kept in a ND free country, zone or compartment from the time they were hatched until the time of slaughter or for at least the 21 days preceding slaughter; or

2. these commodities have been processed to ensure the destruction of NDV (under study);

AND

3. the necessary precautions were taken to avoid contact of the commodity with any source of NDV.

Recommendations for the importation of feathers and down of birds other than poultry

Regardless of the ND status of the country of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities have been processed to ensure the destruction of NDV (under study); and

2. the necessary precautions were taken to avoid contact of the commodity with any source of NDV.

Recommendations for the importation of feather meal

Regardless of the ND status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. these commodities were processed in a ND free country, zone or compartment from poultry which were kept in a ND free country, zone or compartment from the time they were hatched until the time of slaughter or for at least the 21 days preceding slaughter; or

2. these commodities have been processed either:
   a) moist heat at a minimum temperature of 118°C for minimum of 40 minutes; or
   b) with a continuous hydrolysing process under at least 3.79 bar of pressure with steam at a minimum temperature of 122°C for a minimum of 15 minutes;

3. the necessary precautions were taken to avoid contact of the commodity with any source of ND.

Procedures for the inactivation of the ND virus in eggs and egg products

The following times and temperatures are suitable for the inactivation of ND virus present in eggs and egg products:
Temperature (°C) | Time
---|---
Whole egg | 55 | 2521 seconds
Whole egg | 57 | 1596 seconds
Whole egg | 59 | 674 seconds
Liquid egg white | 55 | 2278 seconds
Liquid egg white | 57 | 986 seconds
Liquid egg white | 59 | 301 seconds
10% salted yolk | 59 | 176 seconds
Dried egg white | 57 | 504 hours

The listed temperatures are indicative of a range that achieves a 7-log kill. Where scientifically documented, variances from these times and temperatures may also be suitable when they achieve the inactivation of the virus.

### Article 10.13.19 quadsquint (Under study)

**Procedures for the inactivation of the ND virus in meat**

A procedure which produces a core temperature of 70°C for 574 seconds is suitable for the inactivation of ND virus present in meat.

Temperature (°C) | Time
---|---
Poultry meat | 65.0 | 840 seconds
| 70.0 | 574 seconds
| 74.0 | 280 seconds
| 80.0 | 203 seconds

### Article 10.13.20.

**Surveillance: introduction**

Articles 10.13.20. to 10.13.24. define the principles and provide a guide on the surveillance for ND as defined in Article 10.13.1. and is complementary to Chapter 1.4. It is applicable to Members seeking to determine their ND status. This may be for the entire country, zone or compartment. Guidance for Members seeking free status following an outbreak and for the maintenance of ND status is also provided.

Surveillance for ND is complicated by the known prevalence and occurrence of avian paramyxovirus serotype 1 (APMV-1) infections in many bird species, both domestic and wild, and the widespread utilization of ND vaccines in domestic poultry.
The impact and epidemiology of ND differ widely in different regions of the world and therefore it is not possible to provide specific recommendations for all situations. Therefore, surveillance strategies employed for demonstrating freedom from ND at an acceptable level of confidence will need to be adapted to the local situation. Variables such as the frequency of contacts of poultry with wild birds, different biosecurity levels, production systems and the commingling of different susceptible species require specific surveillance strategies to address each specific situation. It is incumbent upon the Member to provide scientific data that explains the epidemiology of ND in the region concerned and also demonstrates how all the risk factors are managed. There is, therefore, considerable latitude available to Members to provide a well-reasoned argument to prove freedom from NDV infection.

Surveillance for ND should be in the form of a continuing programme designed to establish that the country, zone or compartment, for which application is made, is free from NDV infection.

Article 10.13.21.

**Surveillance: general conditions and methods**

1. A surveillance system in accordance with Chapter 1.4. should be under the responsibility of the Veterinary Authority. In particular there should be in place:

   a) a formal and ongoing system for detecting and investigating outbreaks of disease or NDV infection;

   b) a procedure for the rapid collection and transport of samples from suspect cases of ND to a laboratory for ND diagnosis as described in the Terrestrial Manual;

   c) a system for recording, managing and analysing diagnostic and surveillance data.

2. The ND surveillance programme should:

   a) include an early warning system throughout the production, marketing and processing chain for reporting suspicious cases. Farmers and workers, who have day-to-day contact with poultry, as well as diagnosticians, should report promptly any suspicion of ND to the Veterinary Authority. They should be supported directly or indirectly (e.g. through private veterinarians or veterinary para-professionals) by government information programmes and the Veterinary Authority. All suspected cases of ND should be investigated immediately. As suspicion cannot be resolved by epidemiological and clinical investigation alone, samples should be taken and submitted to a laboratory for appropriate tests. This requires that sampling kits and other equipment are available to those responsible for surveillance. Personnel responsible for surveillance should be able to call for assistance from a team with expertise in ND diagnosis and control;

   b) implement, when relevant, regular and frequent clinical, virological and serological surveillance of high risk groups of poultry within the target population (e.g. those adjacent to an ND infected country, zone, compartment, places where birds and poultry of different origins are mixed, or other sources of NDV).

An effective surveillance system may identify suspicious cases that require follow-up and investigation to confirm or exclude that the cause of the condition is due to NDV infection. The rate at which such suspicious cases are likely to occur will differ between epidemiological situations and cannot therefore be predicted reliably. Applications for freedom from NDV infection should provide details of the occurrence of suspicious cases and how they were investigated and dealt with. This should include the results of laboratory testing and the control measures to which the animals concerned were subjected during the investigation (quarantine, movement stand-still orders, etc.).
Article 10.13.22.

Surveillance strategies

1. Introduction

The principles involved in surveillance for disease/infection are technically well defined. Any surveillance programme requires inputs from professionals competent and experienced in this field and should be thoroughly documented. The design of surveillance programmes to prove the absence of NDV infection/circulation needs to be carefully followed to avoid producing results that are either unreliable, or excessively costly and logistically complicated.

If a Member wishes to declare freedom from NDV infection in a country, zone or compartment, the subpopulation used for surveillance of the disease/infection should be representative of all poultry within the country, zone or compartment. Multiple surveillance methods should be used concurrently to accurately define the true ND status of poultry populations. Active and passive surveillance for ND should be ongoing with the frequency of active surveillance being appropriate to the disease situation in the country. Surveillance should be composed of random and/or targeted approaches, dependent on the local epidemiological situation and using clinical, virological and serological methods as described in the Terrestrial Manual. If alternative tests are used they must have been validated as fit-for-purpose in accordance with OIE standards. A Member should justify the surveillance strategy chosen as adequate to detect the presence of NDV infection in accordance with Chapter 1.4. and the prevailing epidemiological situation.

In surveys, the sample size selected for testing should be statistically justified to detect infection at a predetermined target prevalence. The sample size and expected disease prevalence determine the level of confidence in the results of the survey. The survey design and frequency of sampling should be dependent on the historical and current local epidemiological situation. The Member should justify the choice of survey design and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Chapter 1.4.

Targeted surveillance (e.g. based on the increased likelihood of infection in a population) may be an appropriate strategy.

It may, for example, be appropriate to target clinical surveillance at particular species likely to exhibit clear clinical signs (e.g. unvaccinated chickens). Similarly, virological and serological testing could target species that may not show clinical signs (Article 10.13.2.) of ND and are not routinely vaccinated (e.g. ducks). Surveillance may also target poultry populations at specific risk, for example direct or indirect contact with wild birds, multi-age flocks, local trade patterns including live poultry markets, the presence of more than one species on the holding and poor biosecurity measures in place. In situations where wild birds have been shown to play a role in the local epidemiology of ND, surveillance of wild birds may be of value in alerting Veterinary Services to the possible exposure of poultry, and in particular, of free ranging poultry.
Annex XXII (contd)

The sensitivity and specificity of the diagnostic tests are key factors in the choice of survey design, which should anticipate the occurrence of false positive and false negative reactions. Ideally, the sensitivity and specificity of the tests used should be validated for the vaccination / infection history and for the different species in the target population. If the characteristics of the testing system are known, the rate at which these false reactions are likely to occur can be calculated in advance. There needs to be an effective procedure for following up positives to ultimately determine with a high level of confidence, whether they are indicative of infection or not. This should involve both supplementary tests and follow-up investigation to collect diagnostic material from the original sampling unit as well as flocks which may be epidemiologically linked to it.

The results of active and passive surveillance are important in providing reliable evidence that no NDV infection is present in a country, zone or compartment.

2. Clinical surveillance

Clinical surveillance aims to detect clinical signs suggestive of ND at the flock level and should not be underestimated as an early indication of infection. Monitoring of production parameters (e.g. a drop in feed or water consumption or egg production) is important for the early detection of NDV infection in some populations, as there may be no, or mild clinical signs, particularly if they are vaccinated. Any sampling unit within which suspicious animals are detected should be considered as infected until evidence to the contrary is produced. Identification of infected flocks is vital to the identification of sources of NDV.

A presumptive diagnosis of clinical ND in suspect infected populations should always be confirmed by virological testing in a laboratory. This will enable the molecular, antigenic and other biological characteristics of the virus to be determined.

It is desirable that NDV isolates are sent promptly to an OIE Reference Laboratory for archiving and further characterization if required.

3. Virological surveillance

Virological surveillance should be conducted using tests described in the Terrestrial Manual to:

a) monitor at risk populations;

b) confirm suspect clinical cases;

c) follow up positive serological results in unvaccinated populations or sentinel birds;

d) test ‘normal’ daily mortalities (if warranted by an increased risk e.g. infection in the face of vaccination or in establishments epidemiologically linked to an outbreak).

4. Serological surveillance

Where vaccination is carried out, serological surveillance is of limited value. Serological surveillance cannot be used to discriminate between NDV and other APMV-1. Test procedures and interpretations of results are as described in the Terrestrial Manual. Positive NDV antibody test results can have five possible causes:
a) natural infection with APMV-1;

b) vaccination against ND;

c) exposure to vaccine virus;

d) maternal antibodies derived from a vaccinated or infected parent flock are usually found in the yolk and can persist in progeny for up to 4 weeks;

e) non-specific test reactions.

It may be possible to use serum collected for other survey purposes for ND surveillance. However, the principles of survey design described in these recommendations and the requirement for a statistically valid survey for the presence of NDV should not be compromised.

Discovery of seropositive unvaccinated flocks must be investigated further by conducting a thorough epidemiological investigation. Since seropositive results are not necessarily indicative of infection, virological methods should be used to confirm the presence of NDV in such populations. Until validated strategies and tools to differentiate vaccinated animals from those infected with field APMV-1 are available, serological tools should not be used to identify NDV infection in vaccinated populations.

5. Use of sentinel poultry

There are various applications of the use of sentinel poultry as a surveillance tool to detect virus circulation. They may be used to monitor vaccinated populations or species which are less susceptible to the development of clinical disease for the circulation of virus. Sentinel poultry should be immunologically naïve and may be used in vaccinated flocks. In case of the use of sentinel poultry, the structure and organisation of the poultry sector, the type of vaccine used and local epidemiological factors will determine the type of production systems where sentinels should be placed, the frequency of placement and monitoring of the sentinels.

Sentinel poultry must be in close contact with, but should be identified to be clearly differentiated from, the target population. Sentinel poultry must be observed regularly for evidence of clinical disease and any disease incidents investigated by prompt laboratory testing. The species to be used as sentinels should be proven to be highly susceptible to infection and ideally develop clear signs of clinical disease. Where the sentinel poultry do not necessarily develop overt clinical disease a programme of regular active testing by virological and serological tests should be used (the development of clinical disease may be dependent on the sentinel species used or use of live vaccine in the target population that may infect the sentinel poultry). The testing regime and the interpretation of the results will depend on the type of vaccine used in the target population. Sentinel birds should be used only if no appropriate laboratory procedures are available.

Article 10.13.23.

Documentation of ND free status: additional surveillance procedures

The requirements for a country, zone or compartment to declare freedom from ND are given in Article 10.13.3.
Annex XXII (contd)

A Member declaring freedom of a country, zone or compartment (with or without vaccination) should report the results of a surveillance programme in which the ND susceptible poultry population undergoes regular surveillance planned and implemented according to the general conditions and methods described in these recommendations.

1. **Members declaring freedom from ND for the country, zone or compartment**

   In addition to the general conditions described in the Terrestrial Code, a Member declaring freedom from ND for the entire country, or a zone or a compartment should provide evidence for the existence of an effective surveillance programme. The surveillance programme should be planned and implemented according to general conditions and methods described in this Chapter to demonstrate absence of NDV infection in poultry during the preceding 12 months.

2. **Additional requirements for countries, zones or compartments that practice vaccination**

   Vaccination against ND may be used as a component of a disease prevention and control programme. The vaccine used must comply with the provisions of the Terrestrial Manual.

   In vaccinated populations there is a need to perform surveillance to ensure the absence of NDV circulation. The use of sentinel poultry may provide further confidence of the absence of virus circulation. The surveillance must be repeated at least every 6 months or at shorter intervals according to the risk in the country, zone or compartment, or evidence to show the effectiveness of the vaccination programme is regularly provided.

   **Article 10.13.24.**

**Countries, zones or compartments regaining freedom from ND following an outbreak: additional surveillance procedures**

A Member regaining country, zone or compartment freedom from ND should show evidence of an active surveillance programme depending on the epidemiological circumstances of the outbreak to demonstrate the absence of the infection.

A Member declaring freedom of a country, zone or compartment after an outbreak of ND (with or without vaccination) should report the results of a surveillance programme in which the ND susceptible poultry population undergoes regular surveillance planned and implemented according to the general conditions and methods described in these recommendations.
CHAPTER 11.6.

BOVINE SPONGIFORM ENCEPHALOPATHY

Article 11.6.1.

General provisions and safe commodities

The recommendations in this Chapter are intended to manage the human and animal health risks associated with the presence of the bovine spongiform encephalopathy (BSE) agent in cattle (Bos Taurus and B. indicus) only.

1. When authorising import or transit of the following commodities and any products made from these commodities and containing no other tissues from cattle, Veterinary Authorities should not require any BSE related conditions, regardless of the BSE risk status of the cattle population of the exporting country, zone or compartment:

   a) milk and milk products;

   b) semen and in vivo derived cattle embryos collected and handled in accordance with the recommendations of the International Embryo Transfer Society;

   c) hides and skins;

   d) gelatine and collagen prepared exclusively from hides and skins;

   e) protein-free tallow (maximum level of insoluble impurities of 0.15% in weight) and derivatives made from this tallow;

   f) dicalcium phosphate (with no trace of protein or fat);

   g) deboned skeletal muscle meat (excluding mechanically separated meat) from cattle 30 months of age or less, which were not subjected to a stunning process prior to slaughter, with a device injecting compressed air or gas into the cranial cavity or to a pithing process, and which passed ante-mortem and post-mortem inspections and which has been prepared in a manner to avoid contamination with tissues listed in Article 11.6.14.;

   h) blood and blood by-products, from cattle which were not subjected to a stunning process, prior to slaughter, with a device injecting compressed air or gas into the cranial cavity, or to a pithing process.

2. When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the BSE risk status of the cattle population of the exporting country, zone or compartment.

Standards for diagnostic tests are described in the Terrestrial Manual.
Annex XXIII (contd)

Article 11.6.2.

The BSE risk status of the cattle population of a country, zone or compartment

The BSE risk status of the cattle population of a country, zone or compartment should be determined on the basis of the following criteria:

1. the outcome of a risk assessment, based on the provisions of the Terrestrial Code, identifying all potential factors for BSE occurrence and their historic perspective. Members should review the risk assessment annually to determine whether the situation has changed.

   a) Release assessment

   Release assessment consists of assessing, through consideration of the following, the likelihood that the BSE agent has either been introduced into the country, zone or compartment via commodities potentially contaminated with it, or is already present in the country, zone or compartment:

   i) the presence or absence of the BSE agent in the indigenous ruminant population of the country, zone or compartment and, if present, evidence regarding its prevalence;

   ii) production of meat-and-bone meal or gravers from the indigenous ruminant population;

   iii) imported meat-and-bone meal or gravers;

   iv) imported cattle, sheep and goats;

   v) imported animal feed and feed ingredients;

   vi) imported products of ruminant origin for human consumption, which may have contained tissues listed in Article 11.6.14. and may have been fed to cattle;

   vii) imported products of ruminant origin intended for in vivo use in cattle.

   The results of surveillance and other epidemiological investigations into the disposition of the commodities identified above should be taken into account in carrying out the assessment.

   b) Exposure assessment

   If the release assessment identifies a risk factor, an exposure assessment should be conducted, consisting of assessing the likelihood of cattle being exposed to the BSE agent, through a consideration of the following:

   i) recycling and amplification of the BSE agent through consumption by cattle of meat-and-bone meal or gravers of ruminant origin, or other feed or feed ingredients contaminated with these;

   ii) the use of ruminant carcasses (including from fallen stock), by-products and slaughterhouse waste, the parameters of the rendering processes and the methods of animal feed manufacture;
iii) the feeding or not of ruminants with meat-and-bone meal and greaves derived from ruminants, including measures to prevent cross-contamination of animal feed;

iv) the level of surveillance for BSE conducted on the cattle population up to that time and the results of that surveillance;

2. on-going awareness programme for veterinarians, farmers, and workers involved in transportation, marketing and slaughter of cattle to encourage reporting of all cases showing clinical signs consistent with BSE in target sub-populations as defined in Articles 11.6.20. to 11.6.22.;

3. the compulsory notification and investigation of all cattle showing clinical signs consistent with BSE;

4. the examination carried out in accordance with the Terrestrial Manual in a laboratory of brain or other tissues collected within the framework of the aforementioned surveillance and monitoring system.

When the risk assessment demonstrates negligible risk, the Member should conduct Type B surveillance in accordance with Articles 11.6.20. to 11.6.22.

When the risk assessment fails to demonstrate negligible risk, the Member should conduct Type A surveillance in accordance with Articles 11.6.20. to 11.6.22.

Article 11.6.3.

**Negligible BSE risk**

Commodities from the cattle population of a country, zone or compartment pose a negligible risk of transmitting the BSE agent if the following conditions are met:

1. a risk assessment, as described in point 1 of Article 11.6.2., has been conducted in order to identify the historical and existing risk factors, and the Member has demonstrated that appropriate specific measures have been taken for the relevant period of time defined below to manage each identified risk;

2. the Member has demonstrated that Type B surveillance in accordance with Articles 11.6.20. to 11.6.22. is in place and the relevant points target, in accordance with Table 1, has been met;

3. EITHER:
   a) there has been no case of BSE or, if there has been a case, every case of BSE has been demonstrated to have been imported and has been completely destroyed, and
      i) the criteria in points 2 to 4 of Article 11.6.2. have been complied with for at least 7 years; and
      ii) it has been demonstrated through an appropriate level of control and audit that for at least 8 years neither meat-and-bone meal nor greaves derived from ruminants has been fed to ruminants;

   OR

   b) if there has been an indigenous case, every indigenous case was born more than 11 years ago; and
Annex XXIII (contd)

i) the criteria in points 2 to 4 of Article 11.6.2. have been complied with for at least 7 years; and

ii) it has been demonstrated through an appropriate level of control and audit that for at least 8 years neither meat-and-bone meal nor greaves derived from ruminants has been fed to ruminants; and

iii) all BSE cases, as well as:

- all cattle which, during their first year of life, were reared with the BSE cases during their first year of life, and which investigation showed consumed the same potentially contaminated feed during that period, or

- if the results of the investigation are inconclusive, all cattle born in the same herd as, and within 12 months of the birth of, the BSE cases, if alive in the country, zone or compartment, are permanently identified, and their movements controlled, and, when slaughtered or at death, are completely destroyed.

The Member or zone will be included in the list of negligible risk only after the submitted evidence has been accepted by the OIE. Retention on the list requires that the information for the previous 12 months on surveillance results and feed controls be re-submitted annually and changes in the epidemiological situation or other significant events should be reported to the OIE according to the requirements in Chapter 1.1.

Article 11.6.4.

Controlled BSE risk

Commodities from the cattle population of a country, zone or compartment pose a controlled risk of transmitting the BSE agent if the following conditions are met:

1. a risk assessment, as described in point 1 of Article 11.6.2., has been conducted in order to identify the historical and existing risk factors, and the Member has demonstrated that appropriate measures are being taken to manage all identified risks, but these measures have not been taken for the relevant period of time;

2. the Member has demonstrated that Type A surveillance in accordance with Articles 11.6.20. to 11.6.22. has been carried out and the relevant points target, in accordance with Table 1, has been met; Type B surveillance may replace Type A surveillance once the relevant points target is met;

3. EITHER:

   a) there has been no case of BSE or, if there has been a case, every case of BSE has been demonstrated to have been imported and has been completely destroyed, the criteria in points 2 to 4 of Article 11.6.2. are complied with, and it can be demonstrated through an appropriate level of control and audit that neither meat-and-bone meal nor greaves derived from ruminants has been fed to ruminants, but at least one of the following two conditions applies:

      i) the criteria in points 2 to 4 of Article 11.6.2. have not been complied with for 7 years;

      ii) it cannot be demonstrated that controls over the feeding of meat-and-bone meal or greaves derived from ruminants to ruminants have been in place for 8 years;
b) there has been an indigenous case of BSE, the criteria in points 2 to 4 of Article 11.6.2. are complied with, and it can be demonstrated through an appropriate level of control and audit that neither meat-and-bone meal nor greaves derived from ruminants has been fed to ruminants;

and all BSE cases, as well as:

- all cattle which, during their first year of life, were reared with the BSE cases during their first year of life, and which investigation showed consumed the same potentially contaminated feed during that period, or

- if the results of the investigation are inconclusive, all cattle born in the same herd as, and within 12 months of the birth of, the BSE cases,

if alive in the country, zone or compartment, are permanently identified, and their movements controlled, and, when slaughtered or at death, are completely destroyed.

The Member or zone will be included in the list of controlled risk only after the submitted evidence has been accepted by the OIE. Retention on the list requires that the information for the previous 12 months on surveillance results and feed controls be re-submitted annually and changes in the epidemiological situation or other significant events should be reported to the OIE according to the requirements in Chapter 1.1.

Article 11.6.5.

Undetermined BSE risk

The cattle population of a country, zone or compartment poses an undetermined BSE risk if it cannot be demonstrated that it meets the requirements of another category.

Article 11.6.6.

Recommendations for the importation of bovine commodities from a country, zone or compartment posing a negligible BSE risk

for all commodities from cattle not listed in point 1 of Article 11.6.1.

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the country, zone or compartment complies with the conditions in Article 11.6.3.

Article 11.6.7.

Recommendations for the importation of cattle from a country, zone or compartment posing a negligible BSE risk but where there has been an indigenous case

for cattle selected for export

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:
Annex XXIII (contd)

1. are identified by a permanent identification system in such a way as to demonstrate that they are not exposed cattle as described in point 3b)iii) of Article 11.6.3.;

2. were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants had been effectively enforced.

Article 11.6.8.

Recommendations for the importation of cattle from a country, zone or compartment posing a controlled BSE risk

for cattle

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the country, zone or compartment complies with the conditions referred to in Article 11.6.4.;

2. cattle selected for export are identified by a permanent identification system in such a way as to demonstrate that they are not exposed cattle as described in point 3b) of Article 11.6.4.;

3. cattle selected for export were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants was effectively enforced.

Article 11.6.9.

Recommendations for the importation of cattle from a country, zone or compartment posing an undetermined BSE risk

for cattle

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants has been banned and the ban has been effectively enforced;

2. all BSE cases, as well as:
   a) all cattle which, during their first year of life, were reared with the BSE cases during their first year of life, and, which investigation showed consumed the same potentially contaminated feed during that period, or
   b) if the results of the investigation are inconclusive, all cattle born in the same herd as, and within 12 months of the birth of, the BSE cases, if alive in the country, zone or compartment, are permanently identified, and their movements controlled, and, when slaughtered or at death, are completely destroyed;

3. cattle selected for export:
   a) are identified by a permanent identification system in such a way as to demonstrate that they are not exposed cattle as demonstrated in point 2 above;
   b) were born at least 2 years after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants was effectively enforced.
Annex XXIII (contd)

Article 11.6.10.

Recommendations for the importation of meat and meat products from a country, zone or compartment posing a negligible BSE risk

for fresh meat and meat products from cattle (other than those listed in point 1 of Article 11.6.1.)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the country, zone or compartment complies with the conditions in Article 11.6.3.;
2. the cattle from which the fresh meat and meat products were derived passed ante-mortem and post-mortem inspections;
3. in countries with negligible BSE risk where there have been indigenous cases, the cattle from which the fresh meat and meat products were derived were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and gravers derived from ruminants had been effectively enforced.

Article 11.6.11.

Recommendations for the importation of meat and meat products from a country, zone or compartment posing a controlled BSE risk

for fresh meat and meat products from cattle (other than those listed in point 1 of Article 11.6.1.)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the country, zone or compartment complies with the conditions referred to in Article 11.6.4.;
2. the cattle from which the fresh meat and meat products were derived passed ante-mortem and post-mortem inspections;
3. cattle from which the fresh meat and meat products destined for export were derived were not subjected to a stunning process, prior to slaughter, with a device injecting compressed air or gas into the cranial cavity, or to a pithing process;
4. the fresh meat and meat products were produced and handled in a manner which ensures that such products do not contain and are not contaminated with:
   a) the tissues listed in points 1 and 2 of Article 11.6.14.,
   b) mechanically separated meat from the skull and vertebral column from cattle over 30 months of age.

Article 11.6.12.

Recommendations for the importation of meat and meat products from a country, zone or compartment posing an undetermined BSE risk

for fresh meat and meat products from cattle (other than those listed in point 1 of Article 11.6.1.)
Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the cattle from which the fresh meat and meat products originate:
   a) have not been fed meat-and-bone meal or greaves derived from ruminants;
   b) passed ante-mortem and post-mortem inspections;
   c) were not subjected to a stunning process, prior to slaughter, with a device injecting compressed air or gas into the cranial cavity, or to a pithing process;

2. the fresh meat and meat products were produced and handled in a manner which ensures that such products do not contain and are not contaminated with:
   a) the tissues listed in points 1 and 3 of Article 11.6.14.,
   b) nervous and lymphatic tissues exposed during the deboning process,
   c) mechanically separated meat from the skull and vertebral column from cattle over 12 months of age.

Article 11.6.13.

Recommendations on ruminant-derived meat-and-bone meal or greaves

1. Ruminant-derived meat-and-bone meal or greaves, or any commodities containing such products, which originate from a country, zone or compartment defined in Article 11.6.3., but where there has been an indigenous case of BSE, should not be traded if such products were derived from cattle born before the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants had been effectively enforced.

2. Ruminant-derived meat-and-bone meal or greaves, or any commodities containing such products, which originate from a country, zone or compartment defined in Articles 11.6.4. and 11.6.5. should not be traded between countries.


Recommendations on commodities that should not be traded

1. From cattle of any age originating from a country, zone or compartment defined in Articles 11.6.4. and 11.6.5., the following commodities, and any commodity contaminated by them, should not be traded for the preparation of food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices: tonsils and distal ileum. Protein products, food, feed, fertilisers, cosmetics, pharmaceuticals or medical devices prepared using these commodities (unless covered by other Articles in this Chapter) should also not be traded.

2. From cattle that were at the time of slaughter over 30 months of age originating from a country, zone or compartment defined in Article 11.6.4., the following commodities, and any commodity contaminated by them, should not be traded for the preparation of food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices: brains, eyes, spinal cord, skull and vertebral column. Protein products, food, feed, fertilisers, cosmetics, pharmaceuticals or medical devices prepared using these commodities (unless covered by other Articles in this Chapter) should also not be traded.
3. From cattle that were at the time of slaughter over 12 months of age originating from a country, zone or compartment defined in Article 11.6.5, the following commodities, and any commodity contaminated by them, should not be traded for the preparation of food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices: brains, eyes, spinal cord, skull and vertebral column. Protein products, food, feed, fertilisers, cosmetics, pharmaceuticals or medical devices prepared using these commodities (unless covered by other Articles in this Chapter) should also not be traded.

Article 11.6.15.

**Recommendations for the importation of gelatine and collagen prepared from bones and intended for food or feed, cosmetics, pharmaceuticals including biologicals, or medical devices**

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the commodities came from a country, zone or compartment posing a negligible BSE risk; 

   OR

2. they originate from a country, zone or compartment posing a controlled or undetermined BSE risk and are derived from cattle which have passed ante-mortem and post-mortem inspections; and that
   a) skulls and vertebral columns have been excluded;
   b) the bones have been subjected to a process which includes all of the following steps:
      i) degreasing,
      ii) acid demineralisation,
      iii) acid or alkaline treatment,
      iv) filtration,
      v) sterilisation at >138°C for a minimum of 4 seconds,
      or to an equivalent or better process in terms of infectivity reduction (such as high pressure heating).

Article 11.6.16.

**Recommendations for the importation of tallow (other than as defined in Article 11.6.1) intended for food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices**

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the tallow came from a country, zone or compartment posing a negligible BSE risk; or

2. it originates from a country, zone or compartment posing a controlled BSE risk, is derived from cattle which have passed ante-mortem and post-mortem inspections, and has not been prepared using the tissues listed in points 1 and 2 of Article 11.6.14.
Annex XXIII (contd)

Article 11.6.17.

Recommendations for the importation of dicalcium phosphate (other than as defined in Article 11.6.1.) intended for food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the dicalcium phosphate came from a country, zone or compartment posing a negligible BSE risk; or
2. it originates from a country, zone or compartment posing a controlled or undetermined BSE risk and is a by-product of bone gelatine produced according to Article 11.6.15.

Article 11.6.18.

Recommendations for the importation of tallow derivatives (other than those made from protein-free tallow as defined in Article 11.6.1.) intended for food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the tallow derivatives originate from a country, zone or compartment posing a negligible BSE risk; or
2. they are derived from tallow meeting the conditions referred to in Article 11.6.16.; or
3. they have been produced by hydrolysis, saponification or transesterification using high temperature and pressure.

Article 11.6.19.

Procedures for the reduction of BSE infectivity in meat-and-bone meal

The following procedure should be used to reduce the infectivity of any transmissible spongiform encephalopathy agents which may be present during the production of meat-and-bone meal containing ruminant proteins.

1. The raw material should be reduced to a maximum particle size of 50 mm before heating.
2. The raw material should be heated under saturated steam conditions to a temperature of not less than 133°C for a minimum of 20 minutes at an absolute pressure of 3 bar.

Article 11.6.20.

Surveillance: introduction

1. Depending on the risk category of a country, zone or compartment with regard to bovine spongiform encephalopathy (BSE), surveillance for BSE may have one or more goals:
   a) detecting BSE, to a pre-determined design prevalence, in a country, zone or compartment;
Annex XXIII (contd)

b) monitoring the evolution of BSE in a country, zone or compartment;

c) monitoring the effectiveness of a feed ban and/or other risk mitigation measures, in conjunction with auditing;

d) supporting a claimed BSE status;

e) gaining or regaining a higher BSE status.

2. When the BSE agent is present in a country or zone, the cattle population will comprise the following sectors, in order of decreasing size:

a) cattle not exposed to the infective agent;

b) cattle exposed but not infected;

c) infected cattle, which may lie within one of three stages in the progress of BSE:

i) the majority will die or be killed before reaching a stage at which BSE is detectable by current methods;

ii) some will progress to a stage at which BSE is detectable by testing before clinical signs appear;

iii) the smallest number will show clinical signs.

3. The BSE status of a country, zone or compartment cannot be determined only on the basis of a surveillance programme but should be determined in accordance with all the factors listed in Article 11.6.2. The surveillance programme should take into account the diagnostic limitations associated with the above sectors and the relative distributions of infected cattle among them.

4. With respect to the distribution and expression of the BSE agent within the sectors described above, the following four subpopulations of cattle have been identified for surveillance purposes:

a) cattle over 30 months of age displaying behavioural or clinical signs consistent with BSE (clinical suspects);

b) cattle over 30 months of age that are non-ambulatory, recumbent, unable to rise or to walk without assistance; cattle over 30 months of age sent for emergency slaughter or condemned at ante-mortem inspection (casualty or emergency slaughter or downer cattle);

c) cattle over 30 months of age which are found dead or killed on farm, during transport or at an abattoir (fallen stock);

d) cattle over 36 months of age at routine slaughter.

5. A gradient is used to describe the relative value of surveillance applied to each subpopulation. Surveillance should focus on the first subpopulation, but investigation of other subpopulations will help to provide an accurate assessment of the BSE situation in the country, zone or compartment. This approach is consistent with Articles 11.6.20. to 11.6.22.
Annex XXIII (contd)

6. When establishing a surveillance strategy, authorities need to take into account the inherent difficulties of obtaining samples on farm, and overcome them. These difficulties include higher cost, the necessity to educate and motivate owners, and counteracting potentially negative socio-economic implications.

Article 11.6.21.

Surveillance: description of cattle subpopulations

1. Cattle over 30 months of age displaying behavioural or clinical signs consistent with BSE (clinical suspects)

Cattle affected by illnesses that are refractory to treatment, and displaying progressive behavioural changes such as excitability, persistent kicking when milked, changes in herd hierarchical status, hesitation at doors, gates and barriers, as well as those displaying progressive neurological signs without signs of infectious illness are candidates for examination. These behavioural changes, being very subtle, are best identified by those who handle animals on a daily basis. Since BSE causes no pathognomonic clinical signs, all Members with cattle populations will observe individual animals displaying clinical signs consistent with BSE. It should be recognised that cases may display only some of these signs, which may also vary in severity, and such animals should still be investigated as potential BSE affected animals. The rate at which such suspicious cases are likely to occur will differ among epidemiological situations and cannot therefore be predicted reliably.

This subpopulation is the one exhibiting the highest prevalence. The accurate recognition, reporting and classification of such animals will depend on the ongoing owner/veterinarian awareness programme. This and the quality of the investigation and laboratory examination systems (Article 11.6.2.), implemented by the Veterinary Services, are essential for the credibility of the surveillance system.

2. Cattle over 30 months of age that are non-ambulatory, recumbent, unable to rise or to walk without assistance; cattle over 30 months of age sent for emergency slaughter or condemned at ante-mortem inspection (casualty or emergency slaughter, or downer cattle)

These cattle may have exhibited some of the clinical signs listed above which were not recognised as being consistent with BSE. Experience in Members where BSE has been identified indicates that this subpopulation is the one demonstrating the second highest prevalence. For that reason, it is the second most appropriate population to target in order to detect BSE.

3. Cattle over 30 months of age which are found dead or killed on farm, during transport or at an abattoir (fallen stock)

These cattle may have exhibited some of the clinical signs listed above prior to death, but were not recognised as being consistent with BSE. Experience in Members where BSE has been identified indicates that this subpopulation is the one demonstrating the third highest prevalence.

4. Cattle over 36 months of age at routine slaughter

Experience in Members where BSE has been identified indicates that this subpopulation is the one demonstrating the lowest prevalence. For that reason, it is the least appropriate population to target in order to detect BSE. However, sampling in this subpopulation may be an aide in monitoring the progress of the epizootic and the efficacy of control measures applied, because it offers continuous access to a cattle population of known class, age structure and geographical origin. Testing of routine slaughter cattle 36 months of age or less is of relatively very little value (Table 2).
Surveillance activities

In order to implement efficiently a surveillance strategy for BSE, a Member must use documented records or reliable estimates of the age distribution of the adult cattle population and the number of cattle tested for BSE stratified by age and by subpopulation within the country, zone or compartment.

The approach assigns ‘point values’ to each sample, based on the subpopulation from which it was collected and the likelihood of detecting infected cattle in that subpopulation. The number of points a sample is assigned is determined by the subpopulation from which the sample is collected and the age of the animal sampled. The total points accumulation is then periodically compared to the target number of points for a country, zone or compartment.

A surveillance strategy should be designed to ensure that samples are representative of the herd of the country, zone or compartment, and include consideration of demographic factors such as production type and geographic location, and the potential influence of culturally unique husbandry practices. The approach used and the assumptions made should be fully documented, and the documentation retained for 7 years.

The points targets and surveillance point values in this Chapter were obtained by applying the following factors to a statistical model:

a) the design prevalence for Type A or Type B surveillance;

b) a confidence level of 95%;

c) the pathogenesis, and pathological and clinical expression of BSE:

   i) sensitivity of diagnostic methods used;

   ii) relative frequency of expression by age;

   iii) relative frequency of expression within each subpopulation;

   iv) interval between pathological change and clinical expression;

d) demographics of the cattle population, including age distribution;

e) influence of BSE on culling or attrition of animals from the cattle population via the four subpopulations;

f) percentage of infected animals in the cattle population which are not detected.

Although the procedure accepts very basic information about a cattle population, and can be used with estimates and less precise data, careful collection and documentation of the data significantly enhance their value. Since samples from clinical suspect animals provide many times more information than samples from healthy or dead-of-unknown-cause animals, careful attention to the input data can substantially decrease the procedure’s cost and the number of samples needed. The essential input data are:
Annex XXIII (contd)

g) cattle population numbers stratified by age;

h) the number of cattle tested for BSE stratified by age and by subpopulation.

This Chapter utilises Tables 1 and 2 to determine a desired surveillance points target and the point values of surveillance samples collected.

Within each of the subpopulations above in a country, zone or compartment, a Member may wish to target cattle identifiable as imported from countries or zones not free from BSE and cattle which have consumed potentially contaminated feedstuffs from countries or zones not free from BSE.

All clinical suspects should be investigated, regardless of the number of points accumulated. In addition, animals from the other subpopulations should be tested.

1. Type A surveillance

The application of Type A surveillance will allow the detection of BSE around a design prevalence of at least one case per 100,000 in the adult cattle population in the country, zone or compartment of concern, at a confidence level of 95%.

2. Type B surveillance

The application of Type B surveillance will allow the detection of BSE around a design prevalence of at least one case per 50,000 in the adult cattle population in the country, zone or compartment of concern, at a confidence level of 95%.

Type B surveillance may be carried out by countries, zones or compartments of negligible BSE risk status (Article 11.6.3.) to confirm the conclusions of the risk assessment, for example by demonstrating the effectiveness of the measures mitigating any risk factors identified, through surveillance targeted to maximise the likelihood of identifying failures of such measures.

Type B surveillance may also be carried out by countries, zones or compartments of controlled BSE risk status (Article 11.6.4.), following the achievement of the relevant points target using Type A surveillance, to maintain confidence in the knowledge gained through Type A surveillance.

3. Selecting the points target

The surveillance points target should be selected from Table 1, which shows target points for adult cattle populations of different sizes. The size of the adult cattle population of a country, zone or compartment may be estimated or may be set at one million because, for statistical reasons, one million is the point beyond which sample size does not further increase with population size.
Table 1. Points targets for different adult cattle population sizes in a country, zone or compartment

<table>
<thead>
<tr>
<th>Adult cattle population size (24 months and older)</th>
<th>Type A surveillance</th>
<th>Type B surveillance</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;1,000,000</td>
<td>300,000</td>
<td>150,000</td>
</tr>
<tr>
<td>800,000-1,000,000</td>
<td>240,000</td>
<td>120,000</td>
</tr>
<tr>
<td>600,000-800,000</td>
<td>180,000</td>
<td>90,000</td>
</tr>
<tr>
<td>400,000-600,000</td>
<td>120,000</td>
<td>60,000</td>
</tr>
<tr>
<td>200,000-400,000</td>
<td>60,000</td>
<td>30,000</td>
</tr>
<tr>
<td>100,000-200,000</td>
<td>30,000</td>
<td>15,000</td>
</tr>
<tr>
<td>50,000-100,000</td>
<td>15,000</td>
<td>7,500</td>
</tr>
<tr>
<td>25,000-50,000</td>
<td>7,500</td>
<td>3,750</td>
</tr>
</tbody>
</table>

4. Determining the point values of samples collected

Table 2 can be used to determine the point values of the surveillance samples collected. The approach assigns point values to each sample according to the likelihood of detecting infection based on the subpopulation from which the sample was collected and the age of the animal sampled. This approach takes into account the general principles of surveillance described in Chapter 1.4. and the epidemiology of BSE.

Because precise aging of the animals that are sampled may not be possible, Table 2 combines point values into five age categories. The point estimates for each category were determined as an average for the age range comprising the group. The age groups were selected on their relative likelihoods of expressing BSE according to scientific knowledge of the incubation of the disease and the world BSE experience. Samples may be collected from any combination of subpopulations and ages but should reflect the demographics of the cattle herd of the country, zone or compartment. In addition, Members should sample at least three of the four subpopulations.

Table 2. Surveillance point values for samples collected from animals in the given subpopulation and age category

<table>
<thead>
<tr>
<th>Surveillance subpopulation</th>
<th>Routine slaughter</th>
<th>Fallen stock</th>
<th>Casualty slaughter</th>
<th>Clinical suspect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age=1 year and &lt;2 years</td>
<td>0.01</td>
<td>0.2</td>
<td>0.4</td>
<td>N/A</td>
</tr>
<tr>
<td>Age =2 years and &lt;4 years (young adult)</td>
<td>0.1</td>
<td>0.2</td>
<td>0.4</td>
<td>260</td>
</tr>
<tr>
<td>Age =4 years and &lt;7 years (middle adult)</td>
<td>0.2</td>
<td>0.9</td>
<td>1.6</td>
<td>750</td>
</tr>
<tr>
<td>Age =7 years and &lt;9 years (older adult)</td>
<td>0.1</td>
<td>0.4</td>
<td>0.7</td>
<td>220</td>
</tr>
<tr>
<td>Age =9 years (aged)</td>
<td>0.0</td>
<td>0.1</td>
<td>0.2</td>
<td>45</td>
</tr>
</tbody>
</table>
Annex XXIII (contd)

If a country, zone or compartment determines, based on the demographics and epidemiological characteristics of its cattle population, that precise classification of the subpopulations ‘casualty or emergency slaughter, or downer cattle’ and ‘fallen stock’ is not possible, these subpopulations may be combined. In such a case, the surveillance point values accorded to the combined subpopulation would be that of ‘fallen stock’.

The total points for samples collected may be accumulated over a period of a maximum of 7 consecutive years to achieve the target number of points determined in Table 1.

Surveillance points remain valid for 7 years (the 95th percentile of the incubation period).

Article 11.6.23.

BSE risk assessment: introduction

The first step in determining the BSE risk status of the cattle population of a country or zone is to conduct a risk assessment (reviewed annually), based on Section 2. of this Terrestrial Code, identifying all potential factors for BSE occurrence and their historic perspective.

1. Release assessment

Release assessment consists of assessing the likelihood that a BSE agent has been introduced via the importation of the following commodities potentially contaminated with a BSE agent:

a) meat-and-bone meal or greaves;

b) live animals;

c) animal feed and feed ingredients;

d) products of animal origin for human consumption.

2. Exposure assessment

Exposure assessment consists of assessing the likelihood of exposure of the BSE agent to cattle, through a consideration of the following:

a) epidemiological situation concerning BSE agents in the country or zone;

b) recycling and amplification of the BSE agent through consumption by cattle of meat-and-bone meal or greaves of ruminant origin, or other feed or feed ingredients contaminated with these;

c) the origin and use of ruminant carcasses (including fallen stock), by-products and slaughterhouse waste, the parameters of the rendering processes and the methods of animal feed manufacture;

d) implementation and enforcement of feed bans, including measures to prevent cross-contamination of animal feed.

The following recommendations are intended to assist Veterinary Services in conducting such a risk assessment. They provide guidance on the issues that need to be addressed when conducting a country-based assessment of BSE risk. They apply equally to self-assessment in preparation of dossiers for categorisation of countries. The recommendations are supported by greater detail in the questionnaire used for the submission of data for country assessment.
Article 11.6.24.

The potential for the release of the BSE agent through the importation of meat-and-bone meal or greaves

This point is irrelevant if the exposure assessment outlined below in Article 11.6.27. indicates that meat-and-bone meal or greaves has not been fed, either deliberately or accidentally, in the past 8 years.

Nevertheless, documentation should be provided on the control systems (including relevant legislation) in place to ensure that meat-and-bone meal or greaves has not been fed to ruminants.

Assumption: That meat-and-bone meal or greaves of ruminant origin plays the only significant role in BSE transmission.

Question to be answered: Has meat-and-bone meal, greaves, or feedstuffs containing either been imported within the past 8 years? If so, where from and in what quantities?

Rationale: Knowledge of the origin of meat-and-bone meal, greaves or feedstuffs containing either meat-and-bone meal or greaves, is necessary to assess the risk of release of BSE agent. Meat-and-bone meal and greaves originating in countries of high BSE risk pose a higher release risk than that from low risk countries. Meat-and-bone meal and greaves originating in countries of unknown BSE risk pose an unknown release risk.

Evidence required:

- Documentation to support claims that meat-and-bone meal, greaves or feedstuffs containing either meat-and-bone meal or greaves have not been imported, OR

- Where meat-and-bone meal, greaves or feedstuffs containing them have been imported, documentation of country of origin and, if different, the country of export.

- Documentation on annual volume, by country of origin, of meat, greaves or feedstuffs containing them imported during the past 8 years.

- Documentation describing the composition (on a species and class of stock basis) of the imported meat-and-bone meal, greaves or feedstuffs containing them.

- Documentation, from the country of production, supporting why the rendering processes used to produce meat-and-bone meal, greaves or feedstuffs containing them would have inactivated, or significantly reduced the titre of BSE agent, should it be present.

- Documentation describing the fate of imported meat-and-bone meal and greaves.

Article 11.6.25.

The potential for the release of the BSE agent through the importation of live animals potentially infected with BSE

Assumptions:

- Countries which have imported ruminants from countries infected with BSEs are more likely to experience BSE.
Annex XXIII (contd)

- Cattle pose the only known risk although other species are under study.
- Animals imported for breeding may pose a greater risk than animals imported for slaughter because of the hypothetical risk of maternal transmission and because they are kept to a greater age than animals imported for slaughter.
- Risk is influenced by the date at which imports occurred, relative to the BSE status of the country of origin.
- Risk is proportional to volume of imports (Article 2.2.3.).

Question to be answered: Have live animals been imported within the past 7 years?

Rationale: The release risks are dependent on:

- country of origin and its BSE status, which will change as more data become available; this may result from the detection of clinical disease, or following active surveillance, or assessment of geographical BSE risk;
- feeding and management of the animals in the country of origin;
- use to which the commodity has been put as apart from representing risk of developing clinical disease, the slaughter, rendering and recycling in meat-and-bone meal of imported animals represents a potential route of exposure of indigenous livestock even if meat-and-bone meal and grays, or feedstuffs containing them, have not been imported;
- species;
- dairy versus meat breeds, where there are differences in exposure in the country of origin because feeding practices result in greater exposure of one category;
- age at slaughter.

Evidence required:

- Documentation on the country of origin of imports. This should identify the country of breeding of animals, the length of time they lived in that country and of any other country in which they have resided during their lifetime.
- Documentation describing origins, species and volume of imports.
- Documentation describing the fate of imported animals, including their age at slaughter.
- Documentation demonstrating that risks are periodically reviewed in light of evolving knowledge on the BSE status of the country of origin.

Article 11.6.26.

The potential for the release of the BSE agent through the importation of products of animal origin potentially infected with BSE

Assumptions:

- Semen, embryos, hides and skins or milk are not considered to play a role in the transmission of BSE.
- Countries which have imported products of animal origin from countries with BSEs are more likely to experience BSE.

- Risk is influenced by the date at which imports occurred, relative to the BSE status of the country of origin.

- Risk is proportional to volume of imports (Article 2.2.3).

Question to be answered: What products of animal origin have been imported within the past 7 years?

Rationale The release risks are dependent on:

- the species of origin of the animal products and whether these products contain tissues known to contain BSE infectivity (Article 11.6.14.);

- country of origin and its BSE status, which will change as more data become available; this may result from the detection of clinical disease, or following active surveillance, or assessment of geographical BSE risk;

- feeding and management of the animals in the country of origin;

- use to which the commodity has been put as apart from representing risk of developing clinical disease, the slaughter, rendering and recycling in meat-and-bone meal of imported animals represents a potential route of exposure of indigenous livestock even if meat-and-bone meal and greaves, or feedstuffs containing them, have not been imported;

- species;

- dairy versus meat breeds, where there are differences in exposure in the country of origin because feeding practices result in greater exposure of one category;

- age at slaughter.

Evidence required:

- Documentation on the country of origin of imports. This should identify the country of breeding of animals, the length of time they lived in that country and of any other country in which they have resided during their lifetime.

- Documentation describing origins, species and volume of imports.

- Documentation describing the end use of imported animal products, and the disposal of

- Documentation demonstrating that risks are periodically reviewed in light of evolving knowledge on the BSE status of the country of origin.
Article 11.6.27.

The potential for the exposure of cattle to the BSE agent through consumption of meat-and-bone meal or greaves of ruminant origin

Assumptions:
- That the consumption by bovines of meat-and-bone meal or greaves of ruminant origin plays the only significant role in BSE transmission.
- That commercially-available products of animal origin used in animal feeds may contain meat-and-bone meal or greaves of ruminant origin.
- Milk and blood are not considered to play a role in the transmission of BSE.

Question to be answered: Has meat-and-bone meal or greaves of ruminant origin been fed to cattle within the past 8 years (see Articles 11.6.3. and 11.6.4.)?

Rationale: If cattle have not been fed products of animal origin (other than milk or blood) potentially containing meat-and-bone meal or greaves of ruminant origin within the past 8 years, meat-and-bone meal and greaves can be dismissed as a risk.

Article 11.6.28.

The origin of animal waste, the parameters of the rendering processes and the methods of animal feed production

Assumptions:
- BSE has a long incubation period and insidious onset of signs, so cases may escape detection.
- Pre-clinical BSE infectivity cannot reliably be detected by any method and may enter rendering, in particular if specified risk materials are not removed.
- Tissues most likely to contain high titres of BSE infectivity (brain, spinal cord, eyes) may not be harvested for human consumption and may be rendered.
- BSE may manifest in sudden death, chronic disease, or recumbency, and may be presented as fallen stock or materials condemned as unfit for human consumption.
- BSE agent survival in rendering is affected by the method of processing. Adequate rendering processes are described in Article 11.6.19.
- BSE agent is present at much higher titres in central nervous system and reticulo-endothelial tissues (so-called ‘Specified Risk Materials’, or SRM).

Question to be answered: How has animal waste been processed over the past 8 years?

Rationale: If potentially infected animals or contaminated materials are rendered, there is a risk that the resulting meat-and-bone meal could retain BSE infectivity.
Where meat-and-bone meal is utilized in the production of any animal feeds, the risk of cross-contamination exists.

Evidence required:

- Documentation describing the collection and disposal of fallen stock and materials condemned as unfit for human consumption.
- Documentation describing the definition and disposal of specified risk material, if any.
- Documentation describing the rendering process and parameters used to produce meat-and-bone meal and greaves.
- Documentation describing methods of animal feed production, including details of ingredients used, the extent of use of meat-and-bone meal in any livestock feed, and measures that prevent cross-contamination of cattle feed with ingredients used in monogastric feed.
- Documentation describing monitoring and enforcement of the above.

Article 11.6.29.

Conclusions of the risk assessment

The overall risk of BSE in the cattle population of a country or zone is proportional to the level of known or potential exposure to BSE infectivity and the potential for recycling and amplification of the infectivity through livestock feeding practices. For the risk assessment to conclude that the cattle population of a country or zone is free from BSE risk, it must have demonstrated that appropriate measures have been taken to manage any risks identified.

1 See point 4) of Article 11.6.21.
2 See point 3) of Article 11.6.21.
3 See point 2) of Article 11.6.21.
4 See point 1) of Article 11.6.21.
CHAPTER 11.7.

BOVINE TUBERCULOSIS

Article 11.7.1.

The recommendations in this Chapter are intended to manage the human and animal health risks associated with Mycobacterium \( M. \) \text{tuberculosis complex} (\( M. \) \text{bovis}, \( M. \) \text{caprae}, \( M. \) \text{tuberculosis}, \( M. \) \text{microti}, and \( M. \) \text{africanum}) infection in domestic (permanently captive and owned free-range) bovines including cattle (\text{Bos taurus}, \text{B. indicus} and \text{B. grunniens}), water buffaloes (\text{Bubalus bubalis}) and wood bison (\text{Bison bison} and \text{B. bonasus}).

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 11.7.2.

Country or zone free from bovine tuberculosis

To qualify as free from bovine tuberculosis, a country or zone should satisfy the following requirements:

1. \( M. \) \text{bovis} \text{tuberculosis complex} infection in domestic (permanently captive and owned free-range) bovines including cattle, water buffalo and wood bison is a notifiable disease in the country;
2. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of bovine tuberculosis;
3. regular and periodic testing of all cattle, water buffalo, and wood bison herds did not detect demonstrated that \( M. \) \text{bovis} \text{tuberculosis complex} infection was not present in at least 99.8\% of the herds and 99.9\% of the animals cattle, water buffalo, and wood bison in the country or zone for 3 consecutive years;
4. a surveillance programme should be in place to detect bovine tuberculosis in the country or zone through ante-mortem and post-mortem inspection as described in Chapter 6.2.;
5. if the surveillance programme described in points 3 and 4 above has not detected infection with \( M. \) \text{bovis} \text{tuberculosis complex} for 5 consecutive years, surveillance may be maintained through ante-mortem and post-mortem inspection as described in Chapter 6.2.;
6. cattle, water buffalo and wood bison introduced into a country or zone free from bovine tuberculosis should be accompanied by a certificate from an Official Veterinarian attesting that they come from a country or zone or herd compartment free from bovine tuberculosis or comply with the relevant provisions in Article 11.7.5. or in Article 11.7.6.

Article 11.7.3.

Compartment free from bovine tuberculosis (under study)

To qualify as a compartment free from bovine tuberculosis, a herd or herds of cattle, water buffalo or wood bison in a compartment should be certified by the Veterinary Authority as satisfying the following requirements:
Annex XXIV (contd)

1. cattle, water buffalo and wood bison in the herd or herds:
   a) showed no sign of bovine tuberculosis or lesions at ante-mortem or post-mortem inspection for at least 3 consecutive years;
   b) over 6 weeks of age at the time of the first test and have shown a negative result to at least two tuberculin tests carried out at an interval of a minimum of 6 months, the first test being performed at least 6 months following the slaughter of the last affected animal;
   c) met one of the following conditions:
      i) showed a negative result to an annual biannual tuberculin test to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is more than 1% of all herds in the country or zone during the last 2 years; or
      ii) showed a negative result to an annual tuberculin test every 2 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is more than 0.2% but not more than 1% of all herds in the country or zone during the last 2 years; or
      iii) showed a negative result to a tuberculin test every 3 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is not more than 0.2% of all herds in the country or zone during the last 4 years; or
      iv) showed a negative result to a tuberculin test every 4 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is not more than 0.1% of all herds in the country or zone during the last 6 years;

2. cattle, water buffalo and wood bison introduced into the compartment come from a herd compartment free from bovine tuberculosis. This condition may be waived for animals which have been isolated for at least 90 days and which, prior to entry into the compartment, were subjected to at least two tuberculin tests carried out at a 6-month interval with negative results with the second tuberculin test performed during the 30 days prior to entry into the compartment;

3. cattle, water buffalo and wood bison in a compartment free from bovine tuberculosis are protected from contact with wildlife reservoirs of bovine tuberculosis.

Article 11.7.4.

Herd free from bovine tuberculosis

To qualify as free from bovine tuberculosis, a herd of cattle, water buffalo, or wood bison should satisfy the following requirements:

1. the herd is in a country or a zone free from bovine tuberculosis and is certified free by the Veterinary Authority; or
2. cattle, water buffalo and wood bison in the herd:
   a) showed no signs of bovine tuberculosis or lesions at ante-mortem or post-mortem inspection for at least 3 consecutive years;
   b) over 6 weeks of age, have shown a negative result to at least two tuberculin tests carried out at an interval of a minimum of 6 months, the first test being performed at least 6 months following the slaughter of the last affected animal;
   c) showed a negative result to an annual tuberculin test to ensure the continuing absence of bovine tuberculosis; or
      i) showed a negative result to a tuberculin test every 2 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is not more than 1% of all herds in the country or zone during the last 2 years; or
      ii) showed a negative result to a tuberculin test every 3 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is not more than 0.2% of all herds in the country or zone during the last 4 years; or
      iii) showed a negative result to a tuberculin test every 4 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is not more than 0.1% of all herds in the country or zone during the last 6 years;

3. cattle, water buffalo and wood bison introduced into the herd come from a herd free from bovine tuberculosis. This condition may be waived for animals which have been isolated for at least 90 days and which, prior to entry into the herd, were subjected to at least two tuberculin tests carried out at a 6-month interval with negative results.

Article 11.7.5.

Recommendations for the importation of cattle, water buffalo and wood bison for breeding or rearing

Veterinary authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no signs of bovine tuberculosis on the day of shipment;
2. originate from a herd compartment free from bovine tuberculosis that is in a country, or zone or compartment free from bovine tuberculosis; or
3. were subjected to the tuberculin test for bovine tuberculosis with negative results during the 30 days prior to shipment and come from a herd compartment free from bovine tuberculosis; or
4. have been isolated for at least 90 days prior to entry into the herd including protection from contact with wildlife reservoirs of bovine tuberculosis and were subjected to at least two tuberculin tests carried out at a six-month interval with negative results with the second tuberculin test performed during the 30 days prior to entry into the herd.
Article 11.7.6.

Recommendations for the importation of cattle, water buffalo and wood bison for slaughter

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no signs of bovine tuberculosis on the day of shipment;

2. have been protected from contact with wildlife reservoirs of bovine tuberculosis for 90 days prior to shipment;

3. originated from a herd compartment free from bovine tuberculosis or were subjected to a tuberculin test for bovine tuberculosis with negative results during the 30 days prior to shipment;

4. were not being eliminated as part of an eradication programme against bovine tuberculosis.

Article 11.7.7.

Recommendations for the importation of semen of cattle, water buffalo and wood bison

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no signs of bovine tuberculosis on the day of collection of the semen; and either
   b) were kept in an artificial insemination centre free from bovine tuberculosis in a country, zone or compartment free from bovine tuberculosis and which only accepts animals from a free herd compartment in a free country, zone or compartment; or
   c) showed negative results to tuberculin tests carried out annually and were kept in a herd compartment free from bovine tuberculosis;

2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5.

Article 11.7.8.

Recommendations for the importation of embryos/ova of cattle, water buffalo and wood bison

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) and all other susceptible animals in the herd of origin showed no signs of bovine tuberculosis during the 24 hours prior to embryo collection; and either
   b) originated from a herd compartment free from bovine tuberculosis in a country, or zone or compartment free from bovine tuberculosis; or
were kept in a herd compartment free from bovine tuberculosis, and were subjected to a tuberculin test for bovine tuberculosis with negative results during an isolation period of 30 days in the establishment of origin prior to collection;

2. the embryos/ova were collected, processed and stored in conformity with the provisions of Chapter 4.7., Chapter 4.8. or Chapter 4.9., as relevant.

Article 11.7.9.

Recommendations for the importation of fresh meat and meat products of cattle, water buffalo, and wood bison

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which have been subjected to ante-mortem and post-mortem inspections as described in Chapter 6.2.

Article 11.7.10.

Recommendations for the importation of milk and milk products of cattle, water buffalo and wood bison

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the consignment:

1. has been derived from animals in a herd compartment free from bovine tuberculosis; or
2. was subjected to pasteurization; or
3. was subjected to a combination of control measures with equivalent performance as described in the Codex Alimentarius Code of Hygienic Practice for Milk and Milk Products.
CHAPTER X.X.

BOVINE TUBERCULOSIS OF FARMED CERVIDAE

Article 1.

The recommendations in this Chapter are intended to manage the human and animal health risks associated with Mycobacterium (M.) tuberculosis complex (M. bovis, M. caprae, M. tuberculosis, M. microti and M. africanum) infection in domestic (permanently captive and owned free-range) farmed cervidae (red deer, wapiti, sika, samba, rusa, fallow deer, white-tailed, black-tailed and mule deer [Cervus elephus, C. canadensis, C. nippon, C. unicolor unicolor, C. timorensis, Dama dama dama, Odocoileus virginianus borealis, Odocoileus hemionus columbianus and Odocoileus hemionus hemionus]). The Chapter does not address the management of tuberculosis in wild cervid populations.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 2.

Country or zone free from bovine tuberculosis of farmed cervidae

To qualify as free from bovine tuberculosis of farmed cervidae, a country or zone should satisfy the following requirements:

1. M. bovis tuberculosis complex infection in domestic bovines and in farmed cervidae as specified in Article 1 is a notifiable disease in the country;
2. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of tuberculosis;
3. regular and periodic testing of herds of farmed cervidae has demonstrated that M. bovis tuberculosis complex infection was not present in at least 99.8% of the herds and 99.9% of the farmed cervidae in the country or zone for 3 consecutive years;
4. a surveillance programme should be in place to detect bovine tuberculosis in the country or zone through ante-mortem and post-mortem inspection as described in Chapter 6.2.;
5. if the surveillance programme described in points 3 and 4 above has not detected infection with M. bovis tuberculosis complex in farmed cervidae for 5 consecutive years, surveillance may be maintained through ante-mortem and post-mortem inspection as described in Chapter 6.2.;
6. farmed cervidae introduced into a country or zone free from bovine tuberculosis should be accompanied by a certificate from an Official Veterinarian attesting that they come from a country or zone or compartment free from bovine tuberculosis or comply with the relevant provisions in Article 4 or Article 5.

Article 3.

Compartment free from bovine tuberculosis of farmed cervidae

To qualify as a compartment free from bovine tuberculosis of farmed cervidae, the Veterinary Authority should be able to certify that the following requirements are satisfied:
Annex XXIV (contd)

1. all farmed cervidae:
   a) showed no sign of bovine tuberculosis or lesions at ante-mortem or post-mortem inspection for at least 3 consecutive years;
   b) over 6 weeks of age \textit{at the time of the first test}, have shown a negative result to at least two tuberculin tests carried out at an interval of a minimum of 6 months, the first test being performed at least 6 months following the slaughter of the last affected animal;
   c) met one of the following conditions:
      i) showed a negative result to an annual biannual tuberculin test to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is more than 1\% of all herds in the country or zone during the last 2 years; or
      ii) showed a negative result to an annual tuberculin test every 2 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is more than 0.2\% but not more than 1\% of all herds in the country or zone during the last 2 years; or
      iii) showed a negative result to a tuberculin test every 3 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is not more than 0.2\% of all herds in the country or zone during the last 4 years; or
      iv) showed a negative result to a tuberculin test every 4 years to ensure the continuing absence of bovine tuberculosis if the annual percentage of herds confirmed as infected with tuberculosis is not more than 0.1\% of all herds in the country or zone during the last 6 years;

2. farmed cervidae introduced into the compartment come from a compartment free from bovine tuberculosis. This condition may be waived for animals which have been isolated for at least 90 days and which, prior to entry into the compartment, were subjected to at least two tuberculin tests carried out at a 6-month interval with negative results.

Article 4.

Recommendations for the importation of farmed cervidae for breeding or rearing

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no signs of bovine tuberculosis on the day of shipment;
2. originate from a compartment free from bovine tuberculosis of farmed cervidae that is in a country, or zone, or compartment free from bovine tuberculosis of farmed cervidae; or
3. were subjected to the tuberculin test for bovine tuberculosis with negative results during the 30 days prior to shipment and come from a compartment free from bovine tuberculosis of farmed cervidae; or
4. have been isolated for at least 90 days prior to entry into the herd, including protection from contact with wildlife reservoirs of bovine tuberculosis and were subjected to at least two tuberculin tests carried out at a six-month interval with negative results with the second tuberculin test performed during the 30 days prior to entry into the herd.
Article 5.

Recommendations for the importation of farmed cervidae for slaughter

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no signs of bovine tuberculosis on the day of shipment;
2. have been protected from contact with wildlife reservoirs of bovine tuberculosis for 90 days prior to shipment;
3. originated from a compartment free from bovine tuberculosis of farmed cervidae or were subjected to a tuberculin test for bovine tuberculosis with negative results during the 30 days prior to shipment;
4. were not being eliminated as part of an eradication programme against bovine tuberculosis.

Article 6.

Recommendations for the importation of semen of farmed cervidae

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no signs of bovine tuberculosis on the day of collection of the semen; and either
   b) were kept in an artificial insemination centre compartment free from bovine tuberculosis in any species, in a country, zone or compartment free from bovine tuberculosis of farmed cervidae, and which only accepts animals from a free compartment; or
   c) showed negative results to tuberculin tests carried out annually and were kept in a compartment free from bovine tuberculosis of farmed cervidae;
2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5.

Article 7.

Recommendations for the importation of embryos/ova of farmed cervidae

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) and all other susceptible animals in the herd of origin showed no signs of bovine tuberculosis during the 24 hours prior to embryo collection; and either
   b) originated from a compartment free from bovine tuberculosis of farmed cervidae in a country, zone or compartment free from bovine tuberculosis; or
   c) were kept in a compartment free from bovine tuberculosis of farmed cervidae, and were subjected to a tuberculin test for bovine tuberculosis with negative results during an isolation period of 30 days in the establishment of origin prior to collection;
Annex XXIV (contd)

2. the embryos/ova were collected, processed and stored in conformity with the provisions of Chapter 4.7., Chapter 4.8. or Chapter 4.9., as relevant.

Article 8.

Recommendations for the importation of fresh meat and meat products of farmed cervidae

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which have been subjected to ante-mortem and post-mortem inspections as described in Chapter 6.2.

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CHAPTER 11.8.
CONTAGIOUS BOVINE PLEUROPNEUMONIA

Article 11.8.1.

General provisions

For the purposes of the Terrestrial Code, the incubation period for contagious bovine pleuropneumonia (CBPP) shall be 6 months.

For the purpose of this Chapter, a case of CBPP means an animal infected with Mycoplasma mycoides subsp. mycoides SC (MmmSC), and freedom from CBPP means freedom from MmmSC infection.

For the purpose of this Chapter, susceptible animals include domestic cattle (Bos indicus and B. taurus) and water buffalo (Bubalus bubalis).

For the purposes of international trade, this Chapter deals not only with the occurrence of clinical signs caused by MmmSC, but also with the presence of infection with MmmSC in the absence of clinical signs.

The following defines the occurrence of MmmSC infection:

1. MmmSC has been isolated and identified as such from an animal, embryos, oocytes or semen; or

2. antibodies to MmmSC antigens which are not the consequence of vaccination, or MmmSC DNA, have been identified in one or more animals showing pathological lesions consistent with infection with MmmSC with or without clinical signs, and epidemiological links to a confirmed outbreak of CBPP in susceptible animals.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 11.8.1.bis

Trade in commodities

When authorising import or transit of live ruminants, Veterinary Authorities should comply with recommendations of this Chapter as relevant to the CBPP status of the exporting country, zone or compartment.

When authorising import or transit of the following commodities, Veterinary Authorities should not require any CBPP related conditions, regardless of the CBPP risk status of the bovine domestic cattle and water buffalo population of the exporting country or zone or compartment.
Annex XXV (contd)

1. milk and milk products

2. semen and in vivo derived cattle embryos collected and handled in accordance with the recommendation of the International Embryo Transfer Society,

3. hides and skins;

4. meat and meat products (excluding lung).

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the CBPP status of the domestic cattle and water buffalo population of the exporting country, zone or compartment.

Article 11.8.2.

CBPP free country, zone or compartment

To qualify for inclusion in the existing list of CBPP free countries, a Member should:

1. have a record of regular and prompt animal disease reporting;

2. send a declaration to the OIE stating that:

   a) there has been no outbreak of CBPP during the past 24 months;

   b) no evidence of CBPP infection has been found during the past 24 months;

   c) no vaccination against CBPP has been carried out during the past 24 months,

and supply documented evidence that surveillance for CBPP in accordance with this Chapter is in operation and that regulatory measures for the prevention and control of CBPP have been implemented;

3. not have imported since the cessation of vaccination any animals vaccinated against CBPP.

The country will be included in the list only after the submitted evidence has been accepted by the OIE. Retention on the list requires that the information 2a), 2b), 2c) and 3 above be re-submitted annually and changes in the epidemiological situation or other significant events should be reported to the OIE according to the requirements in Chapter 1.1.

Article 11.8.3.

Recovery of free status

When a CBPP outbreak occurs in a CBPP free country, zone or compartment, one of the following waiting periods is required to regain the status of CBPP free country, zone or compartment:
1. 12 months after the last case where a stamping-out policy and serological surveillance and strict
movement control are applied in accordance with this Chapter;

2. if vaccination was used, 12 months after the slaughter of the last vaccinated animal.

Where a stamping-out policy is not practised, the above waiting periods do not apply but Article 11.8.2.
applies.

Article 11.8.4.

Infected country or zone

When the requirements for acceptance as a CBPP free country, zone or compartment are not fulfilled, a
country shall be considered as CBPP infected.

Article 11.8.5.

Veterinary Authorities of CBPP free countries, zones or compartments may prohibit importation or transit
through their territory of domestic cattle and water buffalo, from countries and zones considered infected
with CBPP.

Article 11.8.6.

Recommendations for importation from CBPP free countries, zones or compartments

for domestic cattle and water buffaloes

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the
animals showed no clinical sign of CBPP on the day of shipment.

Article 11.8.7.

Recommendations for importation from CBPP infected countries or zones

for domestic cattle and water buffaloes for slaughter

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the
animals:

1. showed no clinical sign of CBPP on the day of shipment;

2. originate from an establishment where no case of CBPP was officially reported for the past 6 months, and

3. are transported directly to the slaughterhouse in sealed vehicles.

Article 11.8.7.bis

Recommendations for importation from CBPP free countries, zones or compartments

for bovine semen
V i e t n e r y A u t h o r i t i e s s h o u l d r e q u i r e t h e p r e s e n t a t i o n o f a n i n t e r n a t i o n a l v e t e r i n a r y c e r t i f i c a t e a t t e s t i n g t h a t:

1. t h e d o n o r a n i m a l s:
   a) s h o w e d n o c l i n i c a l s i g n o f C B P P o n t h e d a y o f c o l l e c t i o n o f t h e s e m e n;
   b) w e r e k e e p t i n a C B P P f r e e c o u n t r y s i n c e b i r t h o r f o r a t l e a s t t h e p a s t 6 m o n t h s;

2. t h e s e m e n w a s c o l l e c t e d , p r o c e s s e d a n d s t o r e d i n c o n f o r m i t y w i t h t h e p r o v i s i o n s o f C h a p t e r 4 . 5 .

T e n d i n s e r i a t i o n f r o m C B P P i n f e c t e d c o u n t r i e s o r z o n e s f o r b o v i n e s e m e n

V i e t n e r y A u t h o r i t i e s s h o u l d r e q u i r e t h e p r e s e n t a t i o n o f a n i n t e r n a t i o n a l v e t e r i n a r y c e r t i f i c a t e a t t e s t i n g t h a t:

1. t h e d o n o r a n i m a l s:
   a) s h o w e d n o c l i n i c a l s i g n o f C B P P o n t h e d a y o f c o l l e c t i o n o f t h e s e m e n;
   b) w e r e s u b j e c t e d t o t h e c o m p l e m e n t f i x a t i o n t e s t f o r C B P P w i t h n e g a t i v e r e s u l t s , o n t w o o c c a s i o n s , w i t h a n i n t e r v a l o f n o t l e s s t h a n 2 1 d a y s a n d n o t m o r e t h a n 3 0 d a y s b e t w e e n e a c h t e s t , t h e s e c o n d t e s t b e i n g p e r f o r m e d w i t h i n 1 4 d a y s p r i o r t o c o l l e c t i o n ;
   c) w e r e i s o l a t e d f r o m o t h e r d o m e s t i c b o v i d e a f r o m t h e d a y o f t h e f i r s t c o m p l e m e n t f i x a t i o n t e s t u n t i l c o l l e c t i o n ;
   d) w e r e k e e p e d s i n c e b i r t h , o r f o r t h e p a s t 6 m o n t h s , i n a n e s t a b l i s h m e n t w h e r e n o c a s e o f C B P P w a s r e p o r t e d d u r i n g t h a t p e r i o d , a n d t h a t t h e e s t a b l i s h m e n t w a s n o t s i t u a t e d i n a C B P P i n f e c t e d z o n e .
   e) A N D E I T H E R
      i) h a v e n o t b e e n v a c c i n a t e d a g a i n s t C B P P ;
      OR
      ii) w e r e v a c c i n a t e d u s i n g a v a c c i n e c o m p l i a n t w i t h t h e s t a n d a r d s d e s c r i b e d i n t h e T e r r e s t r i a l M a n u a l n o t m o r e t h a n 4 m o n t h s p r i o r t o c o l l e c t i o n ; i n t h i s c a s e , t h e c o n d i t i o n l a i d d o w n i n p o i n t s b) a b o v e i s n o t r e q u i r e d ;

2. t h e s e m e n w a s c o l l e c t e d , p r o c e s s e d a n d s t o r e d i n c o n f o r m i t y w i t h t h e p r o v i s i o n s o f C h a p t e r 4 . 5 .

O I E T e r r e s t r i a l A n i m a l H e a l t h S t a n d a r d s C o m m i s s i o n / M a r c h 2 0 0 9
Annex XXV (contd)

**Article 11.8.7.quads.**

**Recommendations for importation from CBPP free countries, zones or compartments**

for in vivo derived or in vitro produced embryos/oocytes of bovidae

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of CBPP on the day of collection of the embryos/oocytes;
   b) were kept in a CBPP free country since birth or for at least the past 6 months;

2. the oocytes were fertilised with semen meeting the conditions of Article 11.8.7.bis;

3. the embryos/oocytes were collected, processed and stored in conformity with the provisions of Chapter 4.7., 4.8. or 4.9.

**Article 11.8.7.quints.**

**Recommendations for importation from CBPP infected countries or zone**

for in vivo derived or in vitro produced embryos/oocytes of bovidae

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of CBPP on the day of collection of the embryos/oocytes;
   b) were kept since birth, or for the past 6 months, in an establishment where no case of CBPP was reported during that period, and that the establishment was not situated in a CBPP infected zone;
   c) were isolated from other domestic bovidae from the day of the first complement fixation test until collection;
   d) were vaccinated using a vaccine complying with the standards described in the Terrestrial Manual not more than 4 months prior to collection; in this case, the condition laid down in point b) above is not required;

   **AND EITHER**
   
   i) have not been vaccinated against CBPP;

   **OR**

   ii) were vaccinated using a vaccine complyng with the standards described in the Terrestrial Manual not more than 4 months prior to collection; in this case, the condition laid down in point b) above is not required;
Annex XXV (contd)

2. the oocytes were fertilised with semen meeting the conditions of Article 11.8.7.tris;
3. the embryos/oocytes were collected, processed and stored in conformity with the provisions of Chapter 4.7., 4.8. or 4.9.

Article 11.8.8.

Surveillance: Introduction

The Articles 11.8.9. to 11.8.13. define the principles and provides a guide for the surveillance of contagious bovine pleuropneumonia (CBPP) in accordance with Chapter 1.4. applicable to Members seeking recognition from the OIE for establishment of freedom from CBPP. This may be for the entire country, zone or compartment within the country. Guidance is provided for Members seeking reestablishment of freedom from CBPP for the whole entire country, or for a zone or compartment within the country, following an outbreak, as well as guidelines and for the maintenance of CBPP free status are provided. These guidelines are intended to expand on and explain the requirements of this Chapter. Applications to the OIE for recognition of freedom should follow the format and answer all the questions posed by the “Questionnaire on CBPP” available from the OIE Central Bureau.

The impact and epidemiology of CBPP differ widely in different regions of the world and therefore it is impossible to provide specific recommendations for all situations. It is axiomatic that the Surveillance strategies employed for demonstrating freedom from CBPP at an acceptable level of confidence will need to be adapted to the local situation. It is incumbent upon the applicant Member to submit a dossier to the OIE in support of its application that not only explains the epidemiology of CBPP in the region concerned but also demonstrates how all the risk factors are managed. This should include provision of scientifically-based supporting data. There is therefore considerable latitude available to OIE Members to provide a well-reasoned argument to prove that the absence of CBPP infection is assured at an acceptable level of confidence.

Surveillance for CBPP should be in the form of a continuing programme designed to establish that the whole territory or part of it is free from CBPP infection.

Article 11.8.9.

Surveillance: general conditions and methods

1. A surveillance system in accordance with Chapter 1.4. should be under the responsibility of the Veterinary Authority. A procedure should be in place for the rapid collection and transport of samples from suspect cases of CBPP to a laboratory for CBPP diagnoses as described in the Terrestrial Manual.

2. The CBPP surveillance programme should:

   a) include an early warning system throughout the production, marketing and processing chain for reporting suspicious cases. Farmers and workers (such as community animal health workers) who have day-to-day contact with livestock, meat inspectors as well as laboratory diagnosticians, should report promptly any suspicion of CBPP. They should be integrated directly or indirectly (e.g. through private veterinarians or veterinary para-professionals) into the surveillance system. All suspect cases of CBPP should be investigated immediately. Where suspicion cannot be resolved by epidemiological and clinical investigation, samples should be taken and submitted to a laboratory. This requires that sampling kits and other equipment are available for those responsible for surveillance. Personnel responsible for surveillance should be able to call for assistance from a team with expertise in CBPP diagnosis and control;
b) implement, when relevant, regular and frequent clinical inspection and testing of high-risk groups of animals, such as those adjacent to a CBPP infected country or zone (for example, areas of transhumant production systems);

c) take into consideration additional factors such as animal movement, different production systems, geographical and socio-economic factors that may influence the risk of disease occurrence.

An effective surveillance system will periodically identify suspicious cases that require follow-up and investigation to confirm or exclude that the cause of the condition is CBPP. The rate at which such suspicious cases are likely to occur will differ between epidemiological situations and cannot therefore be predicted reliably. Applications for freedom from CBPP infection should, in consequence, provide details of the occurrence of suspicious cases and how they were investigated and dealt with. This should include the results of laboratory testing and the control measures to which the animals concerned were subjected during the investigation (quarantine, movement stand-still orders, etc.).

Article 11.8.10.

Surveillance strategies

1. Introduction

The target population for surveillance aimed at identifying disease and infection should cover all the susceptible species (Bos taurus, B. indicus and Bubalus bubalis) within the country, zone or compartment to be recognised as free from CBPP infection.

Given the limitations of the diagnostic tools available, the interpretation of surveillance results should be at the herd level rather than at the individual animal level.

Randomised surveillance may not be the preferred approach given the epidemiology of the disease (usually uneven distribution and potential for occult foci of infection in small populations) and the limited sensitivity and specificity of currently available tests. Targeted surveillance (e.g. based on the increased likelihood of infection in particular localities or species, focusing on slaughter findings, and active clinical surveillance) may be the most appropriate strategy. The applicant Member should justify the surveillance strategy chosen as adequate to detect the presence of CBPP infection in accordance with Chapter 1.4. and the epidemiological situation.

Targeted surveillance may involve testing of the entire target subpopulation or a sample from it. In the latter case the sampling strategy will need to incorporate an epidemiologically appropriate design prevalence. The sample size selected for testing will need to be large enough to detect infection if it were to occur at a predetermined minimum rate. The sample size and expected disease prevalence determine the level of confidence in the results of the survey. The applicant Member must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Chapter 1.4. Selection of the design prevalence in particular clearly needs to be based on the prevailing or historical epidemiological situation.
Irrespective of the survey design selected, the sensitivity and specificity of the diagnostic tests employed are key factors in the design, sample size determination and interpretation of the results obtained. Ideally, the sensitivity and specificity of the tests used should be validated.

Irrespective of the surveillance system employed, the design should anticipate the occurrence of false positive reactions. If the characteristics of the testing system are known, the rate at which these false positives are likely to occur can be calculated in advance. There needs to be an effective procedure for following-up positives to ultimately determine with a high level of confidence, whether they are indicative of infection or not. This should involve follow-up with supplementary tests, clinical investigation and post-mortem examination in the original sampling unit as well as herds which may be epidemiologically linked to it.

2. Clinical surveillance

Clinical surveillance aims at detecting clinical signs of CBPP in a herd by close physical examination of susceptible animals. Clinical inspection will be an important component of CBPP surveillance contributing to reach the desired level of confidence of detection of disease if a sufficiently large number of clinically susceptible animals is examined.

Clinical surveillance and laboratory testing should always be applied in series to clarify the status of CBPP suspects detected by either of these complementary diagnostic approaches. Laboratory testing and post-mortem examination may contribute to confirm clinical suspicion, while clinical surveillance may contribute to confirmation of positive serology. Any sampling unit within which suspicious animals are detected should be classified as infected until contrary evidence is produced.

3. Pathological surveillance

Systematic pathological surveillance for CBPP is the most effective approach and should be conducted at slaughterhouses and other slaughter facilities. Suspect pathological findings should be confirmed by agent identification. Training courses for slaughter personnel and meat inspectors are recommended.

4. Serological testing

Serological surveillance is not the preferred strategy for CBPP. However, in the framework of epidemiologic investigations, serological testing may be used.

The limitations of available serological tests for CBPP will make the interpretation of results difficult and useful only at the herd level. Positive findings should be followed-up by clinical and pathological investigations and agent identification.

Clustering of seropositive reactions should be expected in CBPP infections and will be usually accompanied by clinical signs. As clustering may signal field strain infection, the investigation of all instances must be incorporated in the surveillance strategy.

Following the identification of a CBPP infected herd, contact herds need to be tested serologically. Repeated testing may be necessary to reach an acceptable level of confidence in herd classification.
5. **Agent surveillance**

Agent surveillance using tests described in the *Terrestrial Manual* should be conducted to follow-up and confirm or exclude suspect cases. Isolates should be typed to confirm MMMSC.

**Article 11.8.11.**

**Countries or zones applying for recognition of freedom from CBPP**

In addition to the general conditions described in this Chapter, an OIE Member applying for recognition of CBPP freedom for the country or a zone should provide evidence for the existence of an effective surveillance programme. The strategy and design of the surveillance programme will depend on the prevailing epidemiological circumstances and will be planned and implemented according to general conditions and methods in this Chapter, to demonstrate absence of CBPP infection, during the preceding 24 months in susceptible populations. This requires the support of a national or other laboratory able to undertake identification of CBPP infection using methods described in the *Terrestrial Manual*.

**Article 11.8.12.**

**Compartments seeking recognition of freedom from CBPP**

The bilateral recognition of CBPP free compartments should follow the principles laid in this Chapter, Chapter 4.3. and Chapter 4.4.

**Article 11.8.13.**

**Countries or zones re-applying for recognition of freedom from CBPP following an outbreak**

In addition to the general conditions described in this Chapter, a Member re-applying for recognition of country or zone freedom from CBPP should show evidence of an active surveillance programme for CBPP, following the recommendations of this Chapter.

Two strategies are recognised by the OIE in a programme to eradicate CBPP infection following an outbreak:

1. slaughter of all clinically affected and in-contact susceptible animals;
2. vaccination used without subsequent slaughter of vaccinated animals.

The time periods before which an application can be made for re-instatement of freedom from CBPP depends on which of these alternatives is followed. The time periods are prescribed in Article 11.8.3.

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CHAPTER 12.1.

AFRICAN HORSE SICKNESS

Article 12.1.1.

For the purposes of the Terrestrial Code, the infective period for African horse sickness virus (AHSV) shall be 40 days for domestic horses. Although critical information is lacking for some species, this Chapter applies to all equidae.

All countries or zones neighbouring, or considered to be at risk from, a country or zone not having free status should determine their AHSV status from an ongoing surveillance programme. Throughout the Chapter, surveillance is in all cases understood as being conducted as described in Chapter 1.4.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 12.1.2.

AHSV free country or zone

1. A country or zone may be considered free from AHSV when African horse sickness (AHS) is notifiable in the whole country, systematic vaccination is prohibited, importation of equidae and their semen and oocytes or embryos are carried out in accordance with this Chapter, and either:

   a) historical freedom as described in Chapter 1.4. has demonstrated no evidence of AHSV in the country or zone; or

   b) the country or zone has not reported any case of AHS for at least 2 years and is not adjacent to a country or zone not having a free status; or

   c) a surveillance programme has demonstrated no evidence of AHSV in the country or zone for at least 12 months and includes a complete season of vector activity; or

   d) the country or zone has not reported any case of AHS for at least 40 days and a surveillance programme has demonstrated no evidence of Culicoides likely to be competent AHSV vectors for at least 2 years in the country or zone.

2. An AHSV free country or zone will not lose its free status through the importation of vaccinated or seropositive equidae and their semen, oocytes or embryos from infected countries or infected zones, provided these imports are carried out in accordance with this Chapter.

Article 12.1.3.

AHSV seasonally free zone

1. An AHSV seasonally free zone is a part of an infected country or an infected zone for which for part of a year, ongoing surveillance and monitoring consistently demonstrated no evidence of AHSV transmission and no the evidence of the presence of adult Culicoides likely to be competent AHSV vectors.
Annex XXVI (contd)

2. For the application of Articles 12.1.6., 12.1.8. and 12.1.9., the seasonally free period is:
   a) taken to commence the day following the last evidence of AHSV transmission and of the cessation of activity of adult Culicoides likely to be competent AHSV vectors as demonstrated by an ongoing surveillance programme, and
   b) taken to conclude either:
      i) at least 40 days before the earliest date that historical data show AHSV activity has recommenced; or
      ii) immediately when current climatic data or data from a surveillance and monitoring programme indicate an earlier resurgence of activity of adult Culicoides likely to be competent AHSV vectors.

3. An AHSV seasonally free zone will not lose its free status through the importation of vaccinated or seropositive equidae and their semen, oocytes or embryos from infected countries or infected zones, provided these imports are carried out in accordance with this Chapter.

Article 12.1.4.

AHSV infected country or zone

An AHSV infected country or infected zone is one in which the conditions of Article 12.1.2. or Article 12.1.3. do not apply.

Article 12.1.5.

Recommendations for importation from AHSV free countries that are neither neighbouring nor considered to be at risk from an AHSV infected country or infected zone

for equidae

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of AHS on the day of shipment;
2. have not been vaccinated against AHS within the last 40 days;
3. were kept in an AHSV free country since birth or for at least 40 days prior to shipment;
4. either:
   a) did not transit through an infected country or infected zone; or
   b) were protected from attacks by Culicoides at all times when transiting through an infected country or infected zone.
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Article 12.1.6.

Recommendations for importation from AHSV free countries or free zones or from AHSV seasonally free zones (during the seasonally free period) that are neighbouring or are considered to be at risk from an AHSV infected country or infected zone

for equidae

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical signs of AHS on the day of shipment;
2. have not been vaccinated against AHS within the last 40 days;
3. were kept in an AHSV free country, free zone or seasonally free zone during the seasonally free period since birth or for at least 40 days prior to shipment; or
4. in a country or zone considered to be at risk, were held in quarantine for at least 40 days prior to shipment and protected at all times from attacks by Culicoides; and
   a) a serological test according to the Terrestrial Manual to detect antibodies to the AHSV group, was carried out with a negative result on a blood sample collected at least 28 days after introduction into the quarantine station; or
   b) serological tests according to the Terrestrial Manual to detect antibodies against AHSV were carried out with no significant increase in antibody titre on blood samples collected on two occasions, with an interval of not less than 21 days, the first sample being collected at least 7 days after introduction into the quarantine station; or
   c) agent identification tests according to the Terrestrial Manual were carried out with negative results on blood samples collected on two occasions with an interval of not less than 14 days between collection, the first sample being collected at least 7 days after introduction into the quarantine station;
5. were protected from attacks by Culicoides at all times during transportation (including to and at the place of shipment).

Article 12.1.7.

Recommendations for importation from AHSV infected countries or zones

for equidae

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of AHS on the day of shipment;
2. have not been vaccinated against AHS within the last 40 days;
Annex XXVI (contd)

3. were held continuously during the quarantine period of at least 40 days, in a vector-proof quarantine station and protected at all times from attacks by Culicoides; and

   a) a serological test according to the Terrestrial Manual to detect antibodies to the AHSV group, was carried out with a negative result on a blood sample collected at least 28 days after introduction into the quarantine station; or

   b) serological tests according to the Terrestrial Manual to detect antibodies against AHSV were carried out with no significant increase in antibody titre on blood samples collected on two occasions, with an interval of not less than 21 days, the first sample being collected at least 7 days after introduction into the quarantine station; or

   c) agent identification tests according to the Terrestrial Manual were carried out with negative results on blood samples collected on two occasions with an interval of not less than 14 days between collection, the first sample being collected at least 7 days after introduction into the quarantine station;

4. protected from attacks by Culicoides at all times during transportation (including during transportation to and at the place of shipment).

Article 12.1.8.

Recommendations for the importation of equid semen

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the donor animals:

1. showed no clinical sign of AHS on the day of collection of the semen and for the following 40 days;

2. had not been vaccinated against AHS with a live attenuated vaccine within 40 days prior to the day of collection;

3. were either:

   a) kept in an AHSV free country or free zone or from an AHSV seasonally free zone (during the seasonally free period) for at least 40 days before commencement of, and during collection of the semen, or

   b) kept in an AHSV free vector-proof artificial insemination centre throughout the collection period, and subjected to either:

      i) a serological test according to the Terrestrial Manual to detect antibody to the AHSV group, carried out with a negative result on a blood sample collected at least 28 days and not more than 90 days after the last collection of semen; or

      ii) agent identification tests according to the Terrestrial Manual carried out with negative results on blood samples collected at commencement and conclusion of, and at least every 7 days, during semen collection for this consignment.
Article 12.1.9.

Recommendations for the importation of in vivo derived equid embryos/ oocytes

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) showed no clinical sign of AHS on the day of collection of the embryos/oocytes and for the following 40 days;
   b) had not been vaccinated against AHS with a live attenuated vaccine within 40 days prior to the day of collection;
   c) were either:
      i) kept in an AHSV free country or free zone or from an AHSV seasonally free zone (during the seasonally free period) for at least 40 days before commencement of, and during collection of the embryos/oocytes, or
      ii) kept in an AHSV free vector-proof collection centre throughout the collection period, and subjected to either:
         - a serological test according to the Terrestrial Manual to detect antibody to the AHSV group carried out with a negative result on a blood sample collected at least 28 days and not more than 90 days after the last collection of embryos/oocytes; or
         - agent identification tests according to the Terrestrial Manual carried out with negative results on blood samples collected at commencement and conclusion of, and at least every 7 days during embryos/oocytes collection for this consignment;

2. the embryos were collected, processed and stored in conformity with the provisions of Chapter 4.7.;

3. semen used to fertilize the oocytes, complies at least with the requirements in Article 12.1.8.

Article 12.1.10.

Protecting animals from Culicoides attack

When transporting equines through AHSV infected countries or AHSV infected zones, Veterinary Authorities should require strategies to protect animals from attacks by Culicoides during transport, taking into account the local ecology of the vector.

Potential risk management strategies include a combination of:

1. treating animals with chemical repellents prior to and during transportation, in sanitized vehicles treated with appropriate residual contact insecticide;

2. loading, transporting and unloading animals at times of low vector activity (i.e. bright sunshine and low temperature);
3. ensuring vehicles do not stop en route during dawn or dusk, or overnight, unless the animals are held behind insect proof netting;

4. darkening the interior of the vehicle, for example by covering the roof and/or sides of vehicles with shade cloth;

5. monitoring for vectors at common stopping and offloading points to gain information on seasonal variations;

6. using historical, ongoing and/or AHS modelling information to identify low risk ports and transport routes.

Article 12.1.11.

Surveillance: introduction

Articles 12.1.11. to 12.1.13. define the principles and provides a guide on the surveillance for AHS, complementary to Chapter 1.4., applicable to Members seeking to determine their AHSV status. This may be for the entire country or zone. Guidance for Members seeking free status following an outbreak and for the maintenance of AHS status is also provided.

AHS is a vector-borne infection transmitted by a limited number of species of Culicoides insects. Unlike the related bluetongue virus, AHSV is so far geographically restricted to sub Saharan Africa with periodic excursions into North Africa, southwest Europe, the Middle East and adjacent regions of Asia. An important component of AHSV epidemiology is vectorial capacity which provides a measure of disease risk that incorporates vector competence, abundance, seasonal incidence, biting rates, survival rates and the extrinsic incubation period. However, methods and tools for measuring some of these vector factors remain to be developed, particularly in a field context.

According to this Chapter, a Member demonstrating freedom from AHSV infection for the entire country, or a zone should provide evidence for the existence of an effective surveillance programme. The strategy and design of the surveillance programme will depend on the prevailing epidemiological circumstances and should be planned and implemented according to general conditions and methods described in this Chapter. This requires the support of a laboratory able to undertake identification of AHSV infection through the virus detection and antibody tests described in the Terrestrial Manual.

Susceptible wild equid populations should be included in the surveillance programme.

For the purposes of surveillance, a case refers to an equid infected with AHSV.

The purpose of surveillance is to determine if a country or zone is free from AHSV or if a zone is seasonally free from AHSV. Surveillance deals not only with the occurrence of clinical signs caused by AHSV, but also with evidence of infection with AHSV in the absence of clinical signs.

The following defines the occurrence of AHSV infection:

1. AHSV has been isolated and identified as such from an equid or a product derived from that equid,
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2. viral antigen or viral RNA specific to one or more of the serotypes of AHSV has been identified in samples from one or more equids showing clinical signs consistent with AHS, or epidemiologically linked to a confirmed or suspected case, or giving cause for suspicion of previous association or contact with AHSV, or

3. serological evidence of active infection with AHSV by detection of seroconversion with production of antibodies to structural or nonstructural proteins of AHSV that are not a consequence of vaccination have been identified in one or more equids that either show clinical signs consistent with AHS, or epidemiologically linked to a confirmed or suspected case, or give cause for suspicion of previous association or contact with AHSV.

Article 12.1.12.

Surveillance: general conditions and methods

1. A surveillance system should be under the responsibility of the Veterinary Authority. In particular the following should be in place:

a) a formal and ongoing system for detecting and investigating outbreaks of disease;

b) a procedure for the rapid collection and transport of samples from suspect cases of AHS to a laboratory for AHS diagnosis as described in the Terrestrial Manual;

c) a system for recording, managing and analysing diagnostic, epidemiologic and surveillance data.

2. The AHS surveillance programme should:

a) in a country/zone, free or seasonally free, include an early warning system for reporting suspicious cases. Persons who have regular contact with equids, as well as diagnosticians, should report promptly any suspicion of AHS to the Veterinary Authority. An effective surveillance system will periodically identify suspicious cases that require follow-up and investigation to confirm or exclude that the cause of the condition is AHS. The rate at which such suspicious cases are likely to occur will differ between epidemiological situations and cannot therefore be predicted reliably. All suspected cases of AHS should be investigated immediately and samples should be taken and submitted to a laboratory. This requires that sampling kits and other equipment are available for those responsible for surveillance;

b) conduct random or targeted serological and virological surveillance appropriate to the infection status of the country or zone in accordance with Chapter 1.4.

Article 12.1.13.

Surveillance strategies

The target population for surveillance aimed at identification of disease and/or infection should cover susceptible equids within the country or zone. Active and passive surveillance for AHSV infection should be ongoing. Surveillance should be composed of random or targeted approaches using virological, serological and clinical methods appropriate for the infection status of the country or zone.
A Member should justify the surveillance strategy chosen as appropriate to detect the presence of AHSV infection in accordance with Chapter 1.4. and the prevailing epidemiological situation. It may, for example, be appropriate to target clinical surveillance at particular species likely to exhibit clinical signs (e.g. horses). Similarly, virological and serological testing may be targeted to species that rarely show clinical signs (e.g. donkeys).

In vaccinated populations serological and virological surveillance is necessary to detect the AHSV types circulating to ensure that all circulating types are included in the vaccination programme.

If a Member wishes to declare freedom from AHSV infection in a specific zone, the design of the surveillance strategy would need to be aimed at the population within the zone.

For random surveys, the design of the sampling strategy will need to incorporate epidemiologically appropriate design prevalence. The sample size selected for testing will need to be large enough to detect infection if it were to occur at a predetermined minimum rate. The sample size, expected prevalence and diagnostic sensitivity of the tests determine the level of confidence in the results of the survey. The Member must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Chapter 1.4. Selection of the design prevalence, in particular, needs to be based on the prevailing or historical epidemiological situation.

Irrespective of the survey approach selected, the sensitivity and specificity of the diagnostic tests employed are key factors in the design, sample size determination and interpretation of the results obtained. Ideally, the sensitivity and specificity of the tests used should be validated for the vaccination/infection history and the different species in the target population.

Irrespective of the testing system employed, surveillance system design should anticipate the occurrence of false positive reactions. If the characteristics of the testing system are known, the rate at which these false positives are likely to occur can be calculated in advance. There needs to be an effective procedure for following up positives to ultimately determine with a high level of confidence, whether they are indicative of infection or not. This should involve both supplementary tests and follow-up investigation to collect diagnostic material from the original sampling unit as well as those which may be epidemiologically linked to it.

The principles for surveillance for disease/infection are technically well defined. Surveillance programmes to prove the absence of AHSV infection/circulation, need to be carefully designed to avoid producing results that are either insufficiently reliable to be accepted by international trading partners, or excessively costly and logistically complicated. The design of any surveillance programme, therefore, requires inputs from professionals competent and experienced in this field.

1. **Clinical surveillance**

   Clinical surveillance aims at the detection of clinical signs of AHS in equids particularly during a newly introduced infection. In horses, clinical signs may include pyrexia, oedema, hyperaemia of mucosal membranes and dyspnoea.

   AHS suspects detected by clinical surveillance should always be confirmed by laboratory testing.

2. **Serological surveillance**

   Serological surveillance of equid populations is an important tool to confirm absence of AHSV transmission in a country or zone. The species tested should reflect the local epidemiology of AHSV infection, and the equine species available. Management variables that may reduce the likelihood of infection, such as the use of insecticides and animal housing, should be taken into account when selecting equids to be included in the surveillance system.
Samples should be examined for antibodies against AHSV using tests prescribed in the Terrestrial Manual. Positive AHSV antibody tests results can have four possible causes:

a) natural infection with AHSV;

b) vaccination against AHSV;

c) maternal antibodies;

d) positive results due to the lack of specificity of the test.

It may be possible to use sera collected for other purposes for AHSV surveillance. However, the principles of survey design described in these recommendations and the requirements for a statistically valid survey for the presence of AHSV infection should not be compromised.

The results of random or targeted serological surveys are important in providing reliable evidence that no AHSV infection is present in a country or zone. It is, therefore, essential that the survey is thoroughly documented. It is critical to interpret the results in light of the movement history of the animals being sampled.

Serological surveillance in a free zone should target those areas that are at highest risk of AHSV transmission, based on the results of previous surveillance and other information. This will usually be towards the boundaries of the free zone. In view of the epidemiology of AHSV, either random or targeted sampling is suitable to select herds and/or animals for testing.

Serological surveillance in a free country or zone should be carried out over an appropriate distance from the border with an infected country or infected zone, based upon geography, climate, history of infection and other relevant factors. The surveillance should be carried out over a distance of at least 100 kilometres from the border with that country or zone, but a lesser distance could be acceptable if there are relevant ecological or geographical features likely to interrupt the transmission of AHSV. An AHSV free country or zone may be protected from an adjacent infected country or infected zone by a buffer protection zone.

Serological surveillance in infected zones will identify changes in the boundary of the zone, and can also be used to identify the AHSV types circulating. In view of the epidemiology of AHSV infection, either random or targeted sampling is suitable.

3. **Virological surveillance**

Isolation and genetic analysis of AHSV from a proportion of infected animals is beneficial in terms of providing information on serotype and genetic characteristics of the viruses concerned.

Virological surveillance using tests described in the Terrestrial Manual can be conducted:

a) to identify virus circulation in at risk populations;

b) to confirm clinically suspect cases;

c) to follow up positive serological results;

d) to better characterize the genotype of circulating virus in a country or zone.
4. Sentinel animals

Sentinel animals are a form of targeted surveillance with a prospective study design. They comprise groups of unexposed equids that are not vaccinated and are managed at fixed locations and observed and sampled regularly to detect new AHSV infections.

The primary purpose of a sentinel equid programme is to detect AHSV infections occurring at a particular place, for instance sentinel groups may be located on the boundaries of infected zones to detect changes in distribution of AHSV. In addition, sentinel equid programmes allow the timing and dynamics of infections to be observed.

A sentinel equid programme should use animals of known source and history of exposure, control management variables such as use of insecticides and animal housing (depending on the epidemiology of AHSV in the area under consideration), and be flexible in its design in terms of sampling frequency and choice of tests.

Care is necessary in choosing the sites for the sentinel groups. The aim is to maximise the chance of detecting AHSV activity at the geographical location for which the sentinel site acts as a sampling point. The effect of secondary factors that may influence events at each location, such as climate, may also be analysed. To avoid confounding factors sentinel groups should comprise animals selected to be of similar age and susceptibility to AHSV infection. The only feature distinguishing groups of sentinels should be their geographical location. Sera from sentinel animal programmes should be stored methodically in a serum bank to allow retrospective studies to be conducted in the event of new serotypes being isolated.

The frequency of sampling should reflect the equid species used and the reason for choosing the sampling site. In endemic areas virus isolation will allow monitoring of the serotypes and genotypes of AHSV circulating during each time period. The borders between infected and non infected areas can be defined by serological detection of infection. Monthly sampling intervals are frequently used. Sentinel in declared free zones add to confidence that AHSV infections are not occurring unobserved. Here sampling prior to and after the possible period of transmission is sufficient.

Definitive information on AHSV circulating in a country or zone is provided by isolation and identification of the viruses. If virus isolation is required sentinels should be sampled at sufficiently frequent intervals to ensure that some samples are collected during the period of viraemia.

5. Vector surveillance

AHSV is transmitted between equine hosts by species of Culicoides which vary across the world. It is therefore important to be able to identify potential vector species accurately although many such species are closely related and difficult to differentiate with certainty.

The main purpose of vector surveillance is to define high, medium and low-risk areas and local details of seasonality by determining the various species present in an area, their respective seasonal occurrence, and abundance. Vector surveillance has particular relevance to potential areas of spread. Long term surveillance can also be used to assess vector abatement measures.

The most effective way of gathering this information should take account of the biology and behavioural characteristics of the local vector species of Culicoides and may include the use of Onderstepoort-type light traps or similar, operated from dusk to dawn in locations adjacent to equids.
Vector surveillance should be based on scientific sampling techniques. The choice of the number and types of traps to be used in vector surveillance and the frequency of their use should take into account the size and ecological characteristics of the area to be surveyed.

The operation of vector surveillance sites at the same locations as sentinel animals is advisable.

The use of a vector surveillance system to detect the presence of circulating virus is not recommended as a routine procedure as the typically low vector infection rates mean that such detections can be rare. Other surveillance strategies are preferred to detect virus circulation.
CHAPTER 12.7.

EQUINE INFLUENZA

Article 12.7.1.

General provisions

For the purposes of the Terrestrial Code, equine influenza (EI) is defined as an infection of domestic horses, donkeys and mules.

For the purposes of international trade, this Chapter deals not only with the occurrence of clinical signs caused by equine influenza virus (EIV), but also with the presence of infection with EIV in the absence of clinical signs.

For the purposes of this Chapter, isolation is defined as 'the separation of horses from horses of a different equine influenza health status, utilising appropriate biosecurity measures, with the purpose of preventing the transmission of infection'.

For the purposes of the Terrestrial Code, the infective period for equine influenza is 21 days.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 12.7.1.bis

Recommendations on safe Trade in commodities

When authorising import or transit of the following commodities, Veterinary Authorities should not require any EIV related conditions. Regardless of the EI status of the equine population of the exporting country, zone or compartment, the Veterinary Authority of a country, zone or compartment should authorise without restriction on account of EI the importation into their territory of the following commodities:

1. semen;
2. in vivo derived equine embryos collected, processed and stored in conformity with the provisions of Chapter 4.7. (under study).

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the EI status of the equine population of the exporting country, zone or compartment.

Article 12.7.2.

The EI status of a country, a zone or a compartment can be determined on the basis of the following criteria:

1. the outcome of a risk assessment identifying all potential factors for EI occurrence and their historic perspective;
2. whether EI is notifiable in the whole country, an on-going EI awareness programme is in place, and all notified suspect occurrences of EI are subjected to field and, where applicable, laboratory investigations;
3. appropriate surveillance is in place to demonstrate the presence of infection in the absence of clinical signs in horses.
Article 12.7.3.

Equine influenza free country, zone or compartment

A country or a zone or a compartment may be considered free from EI provided the disease is notifiable in the whole country and it shows evidence of an effective surveillance programme, planned and implemented according to the general principles in Chapter 1.4. The surveillance may need to be adapted to parts of the country, zone or compartment depending on historical or geographical factors, industry structure, population data, movements of equids into the country, zone or compartment, wild equid populations or proximity to recent outbreaks.

A country, a zone or a compartment seeking freedom from EI, in which vaccination is practised, should also demonstrate that EIV has not been circulating in the population of domestic and wild horse-equine population during the past 12 months, through surveillance, in accordance with Chapter 1.4. In a country in which vaccination is not practised, surveillance could be conducted using serological testing. In countries where vaccination is practised, the surveillance should include methods of virus detection.

If an outbreak of clinical equine influenza occurs in a previously free country, zone or compartment, free status can be regained 12 months after the last clinical case, providing that surveillance for evidence of infection has been carried out during that 12-month period in accordance with Chapter 1.4.

Article 12.7.4.

Recommendations on safe commodities

Regardless of the EI status of the exporting country, zone or compartment, the Veterinary Authority of a country, zone or compartment should authorise without restriction on account of EI the importation into their territory of the following commodities:

1. semen;
2. in vivo derived equine embryos collected, processed and stored in conformity with the provisions of Chapter 4.7. (under study).

Article 12.7.5.

Recommendations for the importation of horses for immediate slaughter

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the horses showed no clinical sign of EI on the day of shipment.

Article 12.7.6.

Recommendations for the importation of horses for unrestricted movement

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the horses:

1. came from an EI free country, zone or compartment in which they had been resident for at least 21 days; in the case of a vaccinated horse, information on its vaccination status should be included in the veterinary certificate;

OR
2. came from a country, zone or compartment not known to be free from EI, were subjected to pre-export isolation for 21 days and showed no clinical sign of EI during isolation nor on the day of shipment; and

3. were immunised according to the manufacturer's instructions with a vaccine complying with the standards described in the Terrestrial Manual between 21 and 90 days before shipment; and

4. were tested negative for EIV by PCR conducted nasopharyngeal swabs collected on two occasions at 21 days and 3 days before shipment.

Article 12.7.7.

Recommendations for the importation of horses which will be kept in isolation (see Article 12.7.1.)

Veterinary authorities should require the presentation of an international veterinary certificate attesting that the horses:

1. came from an EI free country, zone or compartment in which they had been resident for at least 21 days; in the case of a vaccinated horse, information on its vaccination status should be included in the veterinary certificate;

OR

2. showed no clinical sign of EI in any premises in which the horses had been resident for the 21 days prior to shipment nor on the day of shipment; and

3. were immunised according to the manufacturer's instructions with a vaccine complying with the standards described in the Terrestrial Manual.

Article 12.7.8.

Recommendations for the importation of fresh meat of horses, mules or donkeys

Veterinary authorities should require the presentation of an international veterinary certificate attesting that the fresh meat came from horses, mules or donkeys which had been subjected to ante-mortem and post-mortem inspections as described in Chapter 6.2.
CHAPTER 12.9.

EQUINE RHINOPNEUMONITIS

Article 12.9.1.

General provisions

Equine rhinopneumonitis (ER) is a collective term for any one of several highly contagious, clinical disease entities of equids that may occur as a result of infection by either of two closely related herpesviruses, equid herpesvirus-1 and -4 (EHV-1 and EHV-4).

Infection by either EHV-1 or EHV-4 is characterised by a primary respiratory tract disease of varying severity that is related to the age and immunological status of the infected animal. Infections by EHV-1 in particular are capable of progression beyond the respiratory mucosa to cause the more serious disease manifestations of abortion, perinatal foal death, or neurological dysfunction.

For the purpose of international trade, recommendations are provided for EHV-1 (abortigenic and paralytic forms) only.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 12.9.2.

Recommendations for the importation of equines

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of equine herpes virus type 1 infection (abortigenic and paralytic forms) on the day of shipment and during the 21 days prior to shipment;

2. were kept for the 21 days prior to shipment in an establishment where no case of equine herpes virus type 1 infection (abortigenic and paralytic forms) was reported during that period.
CHAPTER 12.10.

EQUINE VIRAL ARTERITIS

Article 12.10.1.

General provisions

The infective period for equine viral arteritis (EVA) shall be 28 days for all categories of equine except sexually mature stallion where the infective period may be for the life of the animal. Because the infective period may be extended in the case of virus shedding in semen, the status of seropositive stallions should be checked to ensure that they do not shed virus in their semen.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 12.10.2.

Recommendations for the importation of uncastrated male equines

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of EVA on the day of shipment and during the 28 days prior to shipment and met one of the following requirements; and

2. were isolated for the 28 days prior to shipment and were subjected, to a test for EVA, as prescribed in the Terrestrial Manual, carried out either:

   a) on a single blood sample collected during the 28 days prior to shipment with negative result; or

   b) on blood samples taken on two occasions at least 14 days apart within 28 days prior to shipment, which demonstrated stable or declining antibody titres; or

3. were isolated for the 28 days prior to shipment and were subjected between 6 and 9 months of age to a test for EVA, as prescribed in the Terrestrial Manual, carried out on two blood samples collected at least 14 days apart with stable or decreasing titre, immediately vaccinated for EVA and regularly revaccinated according to the manufacturer’s instructions; or

4. were isolated for the 28 days prior to shipment and were subjected to a test for EVA, as prescribed in the Terrestrial Manual, on a blood sample with negative results, immediately vaccinated for EVA, kept for 21 days following vaccination separated from other equidae and regularly revaccinated according to the manufacturer’s instructions; or

   met the following requirements:

   a) were isolated for the 28 days prior to shipment; and

   b) not earlier than 7 days of commencing isolation were tested, with negative results, with a test for EVA as prescribed in the Terrestrial Manual; and

   c) were then immediately vaccinated; and

   d) were kept separated from other equidae for 21 days following vaccination; and
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e) were revaccinated regularly according to the manufacturer’s instructions; or

54. have been subjected to a test for EVA, as prescribed in the Terrestrial Manual, carried out on a blood sample with positive results and then: either

a) were subsequently test mated to two mares within 12 months prior to shipment which were subjected to two tests for EVA as prescribed in the Terrestrial Manual with negative results on blood samples collected at the time of test mating and again 28 days after the mating; or

b) were subjected to a test for equine arteritis virus as prescribed in the Terrestrial Manual with negative results, carried out on semen collected during the 28 days prior to shipment.

Article 12.10.3.

Recommendations for the importation of equines other than uncastrated males

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of EVA on the day of shipment and were kept in an establishment where no animals have shown any signs of EVA for the 28 days prior to shipment; and either

2.1 were isolated for the 28 days prior to shipment and were subjected to a test for EVA, as prescribed in the Terrestrial Manual, carried out either:

a) on a single blood sample collected during the 28 days prior to shipment with negative results, or

b) on blood samples collected on two occasions at least 14 days apart within 28 days prior to shipment, which demonstrated stable or declining antibody titres; or

OR

3.2 were isolated for the 28 days prior to shipment and were subjected, between 6 and 9 months of age, to a diagnostic test for EVA, as prescribed in the Terrestrial Manual on two blood samples collected at least 14 days apart, with negative results or stable or declining titre, and immediately vaccinated for EVA and regularly revaccinated according to the manufacturer’s instructions.

Article 12.10.4.

Recommendations for the importation of semen

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the animal donors:

1. were kept for the 28 days prior to semen collection in an establishment where no equine has shown any clinical sign of EVA during that period; and

2. showed no clinical sign of EVA on the day of semen collection; and

3.1 were subjected between 6 and 9 months of age to a test for EVA as prescribed in the Terrestrial Manual on two blood samples collected at least 14 days apart within 28 days prior to semen collection and which showed a with stable or decreasing titre, immediately vaccinated for EVA and regularly revaccinated according to the manufacturer’s instructions; or
42. were subjected to a test for EVA as prescribed in the Terrestrial Manual on a blood sample with negative results, immediately vaccinated for EVA, kept for 21 days following vaccination separated from other equidae and regularly revaccinated according to the manufacturer’s instructions; or

53. were subjected to a test for EVA as prescribed in the Terrestrial Manual on a blood sample with negative results within 14 days prior to semen collection, and had been separated from other equidae not of an equivalent EVA status for 14 days prior to blood sampling from the time of the taking of the blood sample until the end of semen collection; or

64. have been subjected to a test for EVA as prescribed in the Terrestrial Manual on a blood sample with positive results and then: either

   a) were subsequently test mated to two mares within 12 months prior to semen collection, which were subjected to two tests for EVA as prescribed in the Terrestrial Manual with negative results on blood samples collected at the time of test mating and again 28 days after the test mating, or

   b) were subjected to a test for equine arteritis virus as prescribed in the Terrestrial Manual with negative results, carried out on semen collected within one year prior to collection of the semen to be exported.

75. were, for frozen semen, subjected with negative results either:

   a) to a test for EVA as prescribed in the Terrestrial Manual carried out on a blood sample taken not earlier than 14 days and not later than 12 months after the collection of the semen for export; or

   b) to a test for equine arteritis virus as prescribed in the Terrestrial Manual carried out on an aliquot of the semen collected immediately prior to processing.
CHAPTER 14.9.

SCRAPIE

Article 14.9.1.

General provisions

Scrapie is a neurodegenerative disease of sheep and goats. The main mode of transmission is from mother to offspring immediately after birth and to other susceptible neonates exposed to the birth fluids and tissues of an infected animal. Transmission occurs at a much lower frequency to adults exposed to the birth fluids and tissues of an infected animal. A variation in genetic susceptibility of sheep has been recognised. The incubation period of the disease is variable; however, it is usually measured in years. The duration in incubation period can be influenced by a number of factors including host genetics and strain of agent.

The recommendations in the present Chapter are not intended, or sufficient, to manage the risks associated with the potential presence of the bovine spongiform encephalopathy agent in small ruminants.

Scrapie is not considered to pose a risk to human health. The recommendations in this Chapter are intended to manage the animal health risks associated with the presence of the scrapie agent in sheep and goats. The Chapter does not cover so-called 'atypical' scrapie which is clinically, pathologically, biochemically and epidemiologically unrelated to 'classical' scrapie, may not be contagious and may, in fact, be a spontaneous degenerative condition of older sheep.

1. When authorising import or transit of the following commodities derived from sheep or goats and any products made from these commodities and containing no other tissues from small ruminants derived, Veterinary Authorities should not require any scrapie-related conditions, regardless of the scrapie risk status of the small ruminant sheep and goat populations of the exporting country, zone or compartment:

a) meat (excluding materials as referred to in Article 14.9.11.);

b) semen;

c) hides and skins;

d) gelatine;

e) collagen prepared from hides or skins;

f) tallow (maximum level of insoluble impurities of 0.15% in weight) and derivatives made from this tallow;

g) dicalcium phosphate (with no trace of protein or fat);

h) wool or fibre.

2. When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the scrapie risk status of the small ruminant sheep and goat populations of the exporting country, zone or compartment.

Standards for diagnostic tests are described in the Terrestrial Manual.
Article 14.9.2.

Determination of the scrapie status of the sheep and goat populations of a country, zone or compartment

The scrapie status of the sheep and goat populations of a country, zone or establishment can be determined on the basis of the following criteria:

1. the outcome of a risk assessment identifying all potential factors for scrapie occurrence and their historic perspective, in particular the:
   a) epidemiological situation concerning all animal transmissible spongiform encephalopathies (TSE) in the country, zone or establishment;
   b) importation or introduction of small ruminants sheep and goats or their semen, embryos/oocytes potentially infected with scrapie;
   c) extent of knowledge of the population structure and husbandry practices of sheep and goats in the country or, zone or compartment;
   d) feeding practices, including consumption of meat-and-bone meal or greaves derived from ruminants;
   e) importation of milk and milk products of sheep or goats origin intended for use in feeding of sheep and goats;
   f) importation of meat-and-bone meal or greaves potentially contaminated with an animal TSE or feedstuffs containing either;
   g) the origin and use of ruminant carcasses (including fallen stock), by-products and slaughterhouse waste, the parameters of the rendering processes and the methods of animal feed manufacture;

2. an on-going awareness programme for veterinarians, farmers, and workers involved in transportation, marketing and slaughter of sheep and goats to facilitate recognition and encourage reporting of all animals with clinical signs compatible with scrapie;

3. a surveillance and monitoring system including the following:
   a) official veterinary surveillance, reporting and regulatory control in accordance with the provisions of Chapter 1.4.;
   b) a Veterinary Authority with current knowledge of, and authority over, all establishments which contain sheep and goats in the whole country;
   c) compulsory notification and clinical investigation of all sheep and goats showing clinical signs compatible with scrapie;
   d) examination in accordance with the Terrestrial Manual, in an approved laboratory of appropriate material from sheep and goats older than 18 months displaying clinical signs compatible with scrapie taking into account the recommendations in Chapter X.X. (under study);
   e) maintenance of records including the number and results of all investigations for at least 7 years.
Article 14.9.3.

Scrapie free country or zone

Countries or zones may be considered free from scrapie if within the said territory:

1. a risk assessment, as described in point 1 of Article 14.9.2., has been conducted, and it has been demonstrated that appropriate measures are currently in place and have been taken for the relevant period of time to manage any risk identified and points 2 and 3 have been complied with for the preceding 7 years;

AND EITHER

2. one of the following conditions should be met:
   a) the country or the zone have demonstrated historical freedom taking into account the recommendations in Articles 14.9.13. and 14.9.14.; or

OR

3. for at least 7 years, a surveillance and monitoring system as referred to in Article 14.9.2. has been in place, and no case of scrapie has been reported during this period;

OR

4.b) for at least 7 years, a sufficient number of representative mature culled sheep and goats over 18 months of age investigations have been carried out tested annually, to provide a 95% level of confidence of detecting scrapie if it is present at a prevalence rate exceeding 0.1% out of the total number of all chronic wasting conditions in the population of sheep and goats older than 18 months of age (under study) and no case of scrapie has been reported during this period; it is assumed that the occurrence rate of chronic wasting conditions within the population of sheep and goats older than 18 months of age is at least 1%; or,

OR

5.c) all establishments compartments establishments containing sheep or goats have been accredited free as described in Article 14.9.4.;

AND

6. the feeding to sheep and goats of meat-and-bone meal or greaves potentially contaminated with an animal TSE of ruminant origin has been banned and effectively enforced in the whole country for at least 7 years;

AND

7. introductions of sheep and goats, semen and embryos/oocytes from countries or zones not free from scrapie are carried out in accordance with Articles 14.9.6., 14.9.7., 14.9.8. or 14.9.9., as relevant.

For maintenance of country or zone free status, the investigations referred to in point 4 above should be repeated every 7 years.
Article 14.9.4.

Scrapie free establishment compartment

One or more establishments may be considered eligible for accreditation as a scrapie free establishment compartment if:

1. in the country or zone where the establishment is situated, the following conditions are fulfilled:
   a) the disease is compulsorily notifiable;
   b) an awareness, surveillance and monitoring system as referred to in Article 14.9.2. is in place;
   c) affected sheep and goats are slaughtered and completely destroyed;
   d) the feeding to sheep and goats of meat-and-bone meal or greaves of ruminant origin potentially contaminated with an animal TSE has been banned and effectively enforced in the whole country;
   e) an official accreditation scheme is in operation under the supervision of the Veterinary Authority, including the measures described in point 2 below;

2. in the establishment the following conditions have been complied with for at least 7 years:
   a) sheep and goats should be permanently identified and records maintained, to enable trace back to their establishment of birth;
   b) records of movements of sheep and goats in and out of the establishment are established and maintained;
   c) introductions of animals sheep and goats are allowed only from establishments free country, zone or compartment free establishments of an equal or higher stage in the process of accreditation; however, rams and bucks complying with the provisions in point 2 of Article 14.9.8. may also be introduced;
   d) an Official Veterinarian inspects sheep and goats in the establishment and audits the records at least once a year;
   e) no case of scrapie has been reported;
   f) sheep and goats of the establishment should have no direct or indirect contact, including shared grazing, with sheep or goats from establishments of a lower status;
   g) all culled animals sheep and goats over 18 months of age are inspected by an Official Veterinarian, and a proportion of those exhibiting neurological or wasting signs and all those exhibiting neurological signs are tested in a laboratory for scrapie. The selection of the animals sheep and goats to be tested should be made by the Official Veterinarian. Animals Sheep and goats over 18 months of age that have died or have been killed for reasons other than routine slaughter should also be tested (including ‘fallen’ stock and those sent for emergency slaughter).
Recommendations on safe commodities

Regardless of the scrapie status of the exporting country, Veterinary Authorities should authorise without restriction the import or transit through their territory of meat (excluding materials as referred to in Article 14.9.11.), milk, milk products, wool and its derivatives, hides and skins, tallow, derivatives made from this tallow and dicalcium phosphate originating from sheep and goats.

Recommendations for importation from countries or zones not considered free from scrapie

for sheep and goats for breeding or rearing

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals come from a zone or an establishment compartment free from scrapie as described in Article 14.9.3. and in Article 14.9.4.

OR

In cases where the animals do not come from a compartment free from scrapie as described in Article 14.9.4., the importing country may require the placing of the animals in a quarantine station located on its territory, in conformity with the conditions stipulated in its animal health legislation.

Recommendations for importation from countries or zones not considered free from scrapie

for sheep and goats for slaughter

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. in the country or zone:
   a) the disease is compulsorily notifiable;
   b) an awareness, surveillance and monitoring system as referred to in Article 14.9.2. is in place;
   c) affected sheep and goats are slaughtered and completely destroyed;

2. the sheep and goats selected for export showed no clinical sign of scrapie on the day of shipment.

Recommendations for importation from countries or zones not considered free from scrapie

for semen of sheep and goats

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. in the country or zone:
   a) the disease is compulsorily notifiable;
Annex XXVII (contd)

b) a surveillance and monitoring system as referred to in Article 14.9.2. is in place;

c) affected sheep and goats are slaughtered and completely destroyed;

d) the feeding of sheep and goats with meat and bone meal or greaves potentially contaminated with an animal TSE has been banned and effectively enforced in the whole country;

2. the donor animals:

a) are permanently identified, to enable trace back to their establishment of origin;

b) have been kept since birth in establishments in which no case of scrapie had been confirmed during their residency;

c) showed no clinical sign of scrapie at the time of semen collection;

3. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5.

Article 14.9.8.

Recommendations for importation from countries or zones not considered free from scrapie

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:

a) are permanently identified, to enable trace back to their establishment of origin;

b) have been kept since birth in establishments in which no case of scrapie had been confirmed during their residency;

c) showed no clinical sign of scrapie at the time of semen collection;

2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.5.

Article 14.9.9.

Recommendations for importation from countries or zones not considered free from scrapie

for embryos/oocytes of sheep and goats

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. in the country or zone:

a) the disease is compulsorily notifiable;

b) an awareness, surveillance and monitoring system as referred to in Article 14.9.2. is in place;

c) affected sheep and goats are slaughtered and completely destroyed;
Annex XXVII (contd)

d) the feeding to sheep and goats of meat-and-bone meal or greaves of ruminant origin potentially contaminated with animal TSE has been banned and effectively enforced in the whole country;

2. the donor animals either have been kept since birth in a free compartment, or meet the following conditions:
   a) are permanently identified, to enable trace back to their establishment of origin;
   b) have been kept since birth in establishments in which no case of scrapie had been confirmed during their residency;
   c) showed no clinical sign of scrapie at the time of embryo/oocyte collection;

3. the embryos/oocytes were collected, processed and stored in conformity with the provisions of Chapter 4.7.

Article 14.9.9.bis

Recommendations for importation from countries or zones not considered free from scrapie
for milk and milk products of sheep or goat origin intended for use in feeding of sheep and goats

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the milk and milk products come from scrapie free compartments establishments.

Article 14.9.10.

Recommendations on meat-and-bone meal

Meat-and-bone meal containing any sheep or goat protein, or any feedstuffs containing that type of meat-and-bone meal, which originate from countries not considered free of scrapie should not be traded between countries for ruminant feeding.

Article 14.9.11.

Recommendations for importation from countries or zones not considered free from scrapie
for skulls including brains, ganglia and eyes, vertebral column including ganglia and spinal cord, tonsils, thymus, spleen, intestine, adrenal gland, pancreas, or liver, and protein products derived therefrom, from sheep and goats

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. in the country or zone:
   a) the disease is compulsorily notifiable;
   b) an awareness, surveillance and monitoring system as referred to in Article 14.9.2. is in place;
   c) affected sheep and goats are slaughtered and completely destroyed;

2. the materials come from sheep and goats that showed no clinical sign of scrapie on the day of slaughter.
Annex XXVII (contd)

Article 14.9.12.

**Recommendations for the importation of ovine and caprine materials destined for the preparation of biologicals**

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products originate from sheep and goats born and raised in a scrapie free country, zone or establishment compartment.


**Principles for declaring a country or zone historically free from scrapie**

Articles 14.9.13. and 14.9.14. outline principles for declaring a country or zone free from scrapie.

An essential prerequisite to provide the guarantees required for the recognition of freedom from disease/infection is that the Veterinary Services of the Member comply with the provisions of Chapter 3.1. on evaluation of Veterinary Services, and, if relevant, with the provisions of Chapter 4.3. on zoning and compartmentalisation.

The provisions of the above-mentioned articles are based on the principles developed in Chapter 1.4. and the following premises:

1. the sheep population of the country or zone includes a range of genotypes known to be susceptible to scrapie;
2. the Veterinary Services have the competence, capacity and mandate to investigate, diagnose and report scrapie, if present;
3. the absence of scrapie over a long period of time can be substantiated by effective disease investigation and reporting by the Veterinary Services of an OIE Member.


**Requirements to declare a country or zone historically free from scrapie**

A country or zone may be recognised free from scrapie without having applied the requirements of Article 14.9.3. when:

a) scrapie has been notifiable for at least 25 years; and

b) a formal programme of targeted surveillance and monitoring can be documented as having been in place for at least 10 years; and

c) the presence of a range of scrapie susceptible genotypes in this sheep population can be documented; and

d) appropriate measures to prevent scrapie introduction can be documented as having been in place for at least 25 years; and

i) either scrapie has never been reported; or
ii) no case of scrapie has been reported for at least 25 years.

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CHAPTER 15.3.

CLASSICAL SWINE FEVER

Article 15.3.1.

General provisions

For the purposes of international trade, classical swine fever (CSF) is defined as an infection of domestic pigs.

Domestic pig is defined as ‘all domesticated pigs, permanently captive or farmed free range, used for the production of meat for consumption, for the production of other commercial products or for breeding these categories of pigs.

The pig is the only natural host for classical swine fever (CSF) virus. The definition of pig includes all varieties of Sus scrofa, both domestic breeds and wild boar. For the purposes of this Chapter, a distinction is made between domestic pigs (permanently captive and owned free range pigs) and wild pigs (including feral pigs) populations.

Pigs exposed to CSF virus prenatally may be persistently infected throughout life and may have an incubation period of several months before showing signs of disease. Pigs exposed postnatally have an incubation period of 7-142 days, and are usually infective between post-infection days 5 and 14, but up to 3 months in cases of chronic infections.

For the purposes of international trade, a Member should not impose immediate trade bans in response to a notification of infection with classical swine fever virus in wild pigs according to Article 1.2.3. of the Terrestrial Code.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 15.3.2.

Determination of the CSF status of a country, zone or compartment

The CSF status of a country, zone or compartment can only be determined after considering the following criteria in domestic and wild pigs, as applicable:

1. a risk assessment has been conducted, identifying all potential factors for CSF occurrence and their historic perspective;

2. CSF should be notifiable in the whole territory, and all clinical signs suggestive of CSF should be subjected to appropriate field and/or laboratory investigations;

3. an on-going awareness programme should be in place to encourage reporting of all cases suggestive of CSF;

4. the Veterinary Authority should have current knowledge of, and authority over, all domestic pigs in the country, zone or compartment;

5. the Veterinary Authority should have current knowledge about the population and habitat of wild pigs in the country or zone;
Annex XXVIII (contd)

5. For domestic pigs, appropriate surveillance is in place to capable of detecting the presence of infection even in the absence of clinical signs, and the risk posed by wild pigs is in place; this may be achieved through a surveillance programme in accordance with Articles 15.3.20 to 15.3.26.

6. For wild pigs, if present in the country or zone, a surveillance programme is in place according to Article 15.3.26, taking into account the presence of natural and artificial boundaries, the ecology of the wild pig population, and an assessment of the risks of disease spread, additionally taking measures to limit the spread of CSF within the wild pig population.

7. Based on the assessed risk of spread within the wild pig population, and according to Article 15.3.24, the domestic pig population should be separated from the wild pig population by appropriate biosecurity measures to prevent transmission of CSF from wild to domestic pigs.

Article 15.3.3.

CSF free country, zone or compartment

1. CSF-free status in the absence of an outbreak
   a) Historically free status
      A country or zone or compartment may be considered free from CSF after conducting a risk assessment as referred to in Article 15.3.2, but without formally applying a specific surveillance programme, if the provisions of Article 1.4.6. are complied with.
   b) Free status as a result of a specific surveillance programme
      A country, zone or compartment which does not meet the conditions of point 1 above may be considered free from CSF when a risk assessment as referred to in Article 15.3.2, has been conducted, surveillance in accordance with Articles 15.3.26. to 15.3.31. has been in place for at least 12 months, and when no outbreak has been observed for at least 12 months.
      A country, zone or compartment may be considered free from CSF when surveillance in accordance with Articles 15.3.20. to 15.3.26. has been in place for at least 12 months, and when:

2. CSF-free status following an outbreak
   Free status as a result of an eradication programme
   A country, zone or compartment which does not meet the conditions of point a) or b) above or a compartment may be considered free from CSF if surveillance in accordance with Articles 15.3.26. to 15.3.31. has been in place and after a risk assessment as referred to in Article 15.3.2, has been conducted; and
   a) where a stamping-out policy without vaccination is practised and no outbreak has been observed in domestic pigs for at least 6 months; OR
   b) where a stamping-out policy with vaccination is practised, and either:
      i) vaccinated pigs are slaughtered, and no outbreak has been observed in domestic pigs for at least 6 months after the last vaccinated pig was slaughtered; or
      ii) where there are validated means of distinguishing between vaccinated and infected pigs, no outbreak has been observed in domestic pigs for at least 6 months; OR
c) where a vaccination strategy is practised without a stamping-out policy:
   i) vaccination has been banned in all domestic pigs in the country, zone or compartment for at least 12 months, unless there are validated means of distinguishing between vaccinated and infected pigs;
   ii) if vaccination has been practised within the past 5 years, surveillance in accordance with Articles 15.3.26 to 15.3.31 has been in place for at least 6 months to demonstrate the absence of infection within the population of domestic pigs 6 months to one year old; and
   iii) no outbreak has been observed in domestic pigs for at least 12 months;

AND

in all cases, based on surveillance in accordance with Appendix 3.8.8., CSF infection is not known to occur in any wild pig population in the country or zone

a) there has been no outbreak of CSF in domestic pigs during the past 12 months;

b) no evidence of CSFV infection has been found in domestic pigs during the past 12 months;

c) no vaccination against CSF has been carried out in domestic pigs during the past 12 months;

d) imported domestic pigs comply with the requirements in Article 15.3.5 or Article 15.3.6.

Article 15.3.4.

Country free of CSF in domestic pigs but with a wild pig population

Requirements in points 2a to 2c of Article 15.3.3., as relevant, are complied with. As CSF infection may be present in the wild pig population, the following additional conditions are complied with:

1. a programme for the management of CSF in wild pigs is in place, taking into account the measures in place to manage the disease in the wild pig population, the presence of natural boundaries, the ecology of the wild pig population, and an assessment of the risk of disease spread;

2. zoning or compartmentalisation is applied to prevent transmission of CSF from wild pigs to domestic pigs.

Article 15.3.5.

Recovery of free status

Should a CSF outbreak occur in a previously free country, zone or compartment, the free status of the country, zone or compartment may be restored not less than 30 days after completion of a stamping-out policy where surveillance in accordance with Articles 15.3.26 to 15.3.31 has been carried out with negative results, either:

If emergency vaccination has been practised within the CSF domestic pig control area, recovery of the free status cannot occur before all the vaccinated pigs have been slaughtered, unless there are validated means of distinguishing between vaccinated and infected pigs.

1. 3 months after the last case where a stamping-out policy without vaccination is practised;

OR
Annex XXVIII (contd)

2. where a stamping-out policy with emergency vaccination is practised:
   a) 3 months after the last case and the slaughter of all vaccinated animals; or
   b) 3 months after the last case without the slaughter of vaccinated animals where there are means, validated to OIE standards (Chapter 2.8.3. of the Terrestrial Manual), of distinguishing between vaccinated and infected pigs.

OR

3. where a stamping-out policy is not practised, the provisions of point b)2. of Article 15.3.3. should be followed;

AND

Based on surveillance in accordance with Articles 15.3.20 to 15.3.25., CSFV infection has been demonstrated not to be present in any wild pig population in the country or zone.

Article 15.3.6.

Country or zone free of CSF in wild pigs

A country or zone may be considered free from CSF in wild pigs when:

1. the domestic pig population in the country or zone is free from CSF infection;

2. surveillance in accordance with Articles 15.3.26. to 15.3.31. has been in place to determine the CSF status of the wild pig population in the country, and in the country or zone
   a) there has been no clinical evidence, nor virological evidence of CSF in wild pigs during the past 12 months;
   b) no seropositive wild pigs have been detected in the age class 6-12 months during the past 12 months;

3. there has been no vaccination in wild pigs for the past 12 months;

4. the feeding of swill to wild pigs is forbidden, unless the swill has been treated to destroy any CSF virus that may be present, in conformity with one of the procedures referred to in Article 15.3.24.;

5. imported wild pigs comply with the relevant requirements set forth in the present chapter.

Article 15.3.25.

Recommendations for importation from countries, zones or compartments free of CSF

for domestic pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of CSF on the day of shipment;

2. were kept in a country, zone or compartment free of CSF since birth or for at least the past 3 months;
3. have not been vaccinated against CSF, nor are they the progeny of vaccinated sows, unless there are means, validated to OIE standards (Chapter 2.8.3. of the Terrestrial Manual), of distinguishing between vaccinated and infected pigs.

**Article 15.3.8.**

**Recommendations for importation from countries free of CSF in domestic pigs but with a wild pig population**

*for domestic pigs*

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. were kept in a country or zone free of CSF in domestic pigs since birth or for at least the past 3 months;
2. have not been vaccinated against CSF, nor are they the progeny of vaccinated sows, unless there are validated means of distinguishing between vaccinated and infected pigs;
3. come from a CSF free zone or compartment;
4. showed no clinical sign of CSF on the day of shipment.

**Article 15.3.9.**

**Recommendations for importation from CSF infected countries or zones with CSF infection in domestic pigs**

*for domestic pigs*

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. have not been vaccinated against CSF nor are they the progeny of vaccinated sows, unless there are validated means of distinguishing between vaccinated and infected pigs showed no clinical sign of CSF on the day of shipment;
2. were kept since birth or for the past 3 months in a CSF free compartment;
3. showed no clinical sign of CSF on the day of shipment have not been vaccinated against CSF nor are they the progeny of vaccinated sows, unless there are means, validated to OIE standards (Chapter 2.8.3. of the Terrestrial Manual), of distinguishing between vaccinated and infected pigs.

**Article 15.3.10.**

**Recommendations for importation from countries or zones free of CSF of wild pigs**

*for wild pigs*

Regardless of the CSF status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of CSF on the day of shipment;
Annex XXVIII (contd)

2. have been captured in a country or zone free from CSF were kept in a quarantine station for 40 days prior to shipment, and were subjected to a virological test and a serological test performed at least 21 days after entry into the quarantine station, with negative results.

3. have not been vaccinated against CSF, unless there are validate means, validated to OIE standards (Chapter 2.8.3. of the Terrestrial Manual), of distinguishing between vaccinated and infected pigs and, if the zone where the animal has been captured is adjacent to a zone with infection in wild pigs:

4. were kept in a quarantine station for 40 days prior to shipment, and were subjected to a virological test and a serological test performed at least 21 days after entry into the quarantine station, with negative results.

Article 15.3.118.

Recommendations for importation from countries, zones or compartments free of CSF

for semen of domestic pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) were kept in a country, zone or compartment free of CSF since birth or for at least 3 months prior to collection;
   b) showed no clinical sign of CSF on the day of collection of the semen;

2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.6.

Article 15.3.12.

Recommendations for importation from countries free of CSF in domestic pigs but with a wild pig population

for semen of domestic pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) were kept in a country, zone or compartment free of CSF in domestic pigs since birth or for at least 3 months prior to collection;
   b) showed no clinical sign of CSF on the day of collection of the semen and for the following 40 days;

2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.6.

Article 15.3.139.

Recommendations for importation from CSF infected countries or zones considered infected with CSF in domestic pigs
for semen of domestic pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) were kept in a compartment free of CSF in domestic pigs since birth or for at least 3 months prior to collection;
   b) showed no clinical sign of CSF on the day of collection of the semen and for the following 40 days;
   c) met one of the following conditions:
      i) have not been vaccinated against CSF and were subjected to a serological test performed at least 21 days after collection, with negative results;
      ii) have been vaccinated against CSF and were subjected to a serological test in accordance with the Terrestrial Manual performed at least 21 days after collection and it has been conclusively demonstrated by means validated to OIE standards (Chapter 2.8.3. of the Terrestrial Manual) that any antibody is due to the vaccine; or
      iii) have been vaccinated against CSF and were subjected to a virological test performed in accordance with the Terrestrial Manual on a sample taken on the day of collection and it has been conclusively demonstrated that the boar is negative for virus genome;

2. the semen was collected, processed and stored in conformity with the provisions of Chapter 4.6.

Article 15.3.1410.

Recommendations for importation from countries, zones or compartments free of CSF
for in vivo derived embryos of domestic pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor females showed no clinical sign of CSF on the day of collection of the embryos;

2. the embryos were collected, processed and stored in conformity with the provisions of Chapter 4.7.

Article 15.3.15.

Recommendations for importation from countries free of CSF in domestic pigs but with a wild pig population
for in vivo derived embryos of pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) were kept in a country, zone or compartment free of CSF in domestic pigs since birth or for at least 3 months prior to collection;
Annex XXVIII (contd)

b) showed no clinical sign of CSF on the day of collection of the embryos;

2. the embryos were collected, processed and stored in conformity with the provisions of Chapter 4.7.

Article 15.3.16

Recommendations for importation from **CSF infected** countries or zones **considered infected with CSF in domestic pigs**

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) were kept in a CSF free compartment in domestic pigs since birth or for at least 3 months prior to collection;
   b) showed no clinical sign of CSF on the day of collection of the embryos and for the following 40 days;
   c) and either
      i) have not been vaccinated against CSF and were subjected, with negative results, to a serological test performed at least 21 days after collection; or
      ii) have been vaccinated against CSF and were subjected to a serological test performed at least 21 days after collection and it has been conclusively demonstrated by means, validated to OIE standards (Chapter 2.8.3. of the Terrestrial Manual), that any antibody is due to the vaccine;

2. the embryos were collected, processed and stored in conformity with the provisions of Chapter 4.7.

Article 15.3.17

Recommendations for importation from countries, zones or compartments free of CSF

for **fresh meat of domestic pigs**

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which:

1. have been kept in a country, zone or compartment free of CSF since birth or for at least the past 3 months, or which have been imported in accordance with Article 15.3.5. or Article 15.3.6;

2. have been slaughtered in an approved abattoir, have been subjected to ante-mortem and post-mortem inspections in accordance with Chapter 6.2, and have been found free of any sign suggestive of CSF.

Article 15.3.18

Recommendations for importation from countries or zones free of CSF in domestic pigs but with a wild pig population
Annex XXVIII (contd)

for fresh meat of domestic pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which:

1. were kept in a country, zone or compartment free of CSF in domestic pigs since birth or for at least the past 3 months;

2. have been slaughtered in an approved abattoir, have been subjected to ante-mortem and post-mortem inspections as described in the Codex Alimentarius Code of Hygienic Practice for Meat and have been found free of any sign suggestive of CSF.

Article 15.3.19.

Recommendations for importation from countries or zones free of CSF of fresh meat of wild pigs

Regardless of the CSF status of the country, zone or compartment of origin, Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the entire consignment of meat comes from animals which:
   a) have been killed in a CSF free country or zone;
   b) which have been subjected to a post-mortem inspection as described in the Codex Alimentarius Code of Hygienic Practice for Meat in accordance with Chapter 6.2, in an approved examination centre, and have been found free of any sign suggestive of CSF;

2. from each of which a sample has been collected and has been subjected to a virological test and a serological test for CSF, with negative results.

2. a sample has been collected from every animal shot killed, and has been subjected to a virological test and a serological test for CSF, with negative results.

Article 15.3.20.

Recommendations for the importation of meat and meat products of pigs (either domestic or wild), or for products of animal origin (from fresh meat of pigs) intended for use in animal feeding, for agricultural or industrial use, or for pharmaceutical or surgical use, or for trophies derived from wild pigs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. have been prepared:
   a) exclusively from fresh meat meeting the conditions laid down in Articles 15.3.17.12, 15.3.18, or 15.3.19, as relevant;
   b) in a processing establishment:
      i) approved by the Veterinary Authority for export purposes;
      ii) processing only meat meeting the conditions laid down in Articles 15.3.17.12, 15.3.18, or 15.3.19, as relevant;
OR

2. have been processed in an establishment approved by the Veterinary Authority for export purposes so as to ensure the destruction of the CSF virus in conformity with one of the procedures referred to in Article 15.3.2519, and that the necessary precautions were taken after processing to avoid contact of the product with any source of CSF virus.

Recommendations for the importation of products of animal origin (from pigs, but not derived from fresh meat) intended for use in animal feeding and for agricultural or industrial use

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. originated from domestic pigs in a CSF free country, zone or compartment and have been prepared:
   a) exclusively from products meeting the conditions laid down for fresh meat in Articles 15.3.1712, 15.3.18, or 15.3.1913, as relevant;
   b) in a processing establishment:
      i) approved by the Veterinary Authority for export purposes; or
      ii) processing only products meeting the conditions laid down in point a) above.

OR

2. have been processed in an establishment approved by the Veterinary Authority for export purposes so as to ensure the destruction of the CSF virus in conformity with one of the procedures referred to in Article 15.3.2519, and that the necessary precautions were taken after processing to avoid contact of the product with any source of CSF virus.

Recommendations for the importation of bristles (from pigs)

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:
1. come from originated from domestic pigs in a CSF free country, zone or compartment; or and have been prepared in a processing establishment approved by the Veterinary Authority for export purposes; or

2. have been processed in an establishment approved by the Veterinary Authority for export purposes so as to ensure the destruction of the CSF virus (under study) and that the necessary precautions were taken after processing to avoid contact of the product with any source of CSF virus.

Article 15.3.23

Recommendations for the importation of litter and manure (from pigs)

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. come from originate from domestic pigs in a country, zone or compartment free of CSF; or and have been prepared in a processing establishment approved by the Veterinary Authority for export purposes; or

2. have been processed in an establishment approved by the Veterinary Authority for export purposes so as to ensure the destruction of the CSF virus (under study) and that the necessary precautions were taken after processing to avoid contact of the product with any source of CSF virus.

Article 15.3.17bis

Recommendations for the importation of skins and trophies derived from wild pigs

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the products:

1. originate from domestic pigs in a country, zone or compartment free of CSF and have been prepared in a processing establishment approved by the Veterinary Authority for export purposes; or

2. have been processed in an establishment approved by the Veterinary Authority for export purposes so as to ensure the destruction of the CSF virus in conformity with one of the procedures referred to in Article 15.3.19bis and that the necessary precautions were taken after processing to avoid contact of the product with any source of CSF virus.

Article 15.3.2418

Procedures for the inactivation of the CSF virus in swill

For the inactivation of CSF viruses likely to be present in swill, one of the following procedures should be used:

1. the swill should be maintained at a temperature of at least 90°C for at least 60 minutes, with continuous stirring; or

2. the swill should be maintained at a temperature of at least 121°C for at least 10 minutes at an absolute pressure of 3 bar.

Article 15.3.2519

Procedures for the inactivation of the CSF virus in meat

OIE Terrestrial Animal Health Standards Commission / March 2009
Annex XXVIII (contd)

For the inactivation of viruses present in meat, one of the following procedures should be used:

1. **Heat treatment**
   
   Meat shall be subjected to one of the following treatments:
   
   a) heat treatment in a hermetically sealed container with a Fo value of 3.00 or more;
   
   b) heat treatment at a minimum temperature of 70°C, which must be reached throughout the meat.

2. **Natural fermentation and maturation**
   
   The meat should be subjected to a treatment consisting of natural fermentation and maturation having the following characteristics:
   
   a) an aw value of not more than 0.93, or
   
   b) a pH value of not more than 6.0.

   Hams should be subjected to a natural fermentation and maturation process for at least 190 days and loins for 140 days.

3. **Dry cured pork meat**
   
   a) Italian style hams with bone-in should be cured with salt and dried for a minimum of 313 days.
   
   b) Spanish style pork meat with bone-in should be cured with salt and dried for a minimum of 252 days for Iberian hams, 140 days for Iberian shoulders, 126 days for Iberian loin, and 140 days for Serrano hams.

**Article 15.3.19.bis**

**Procedures for the inactivation of the CSF virus in trophies**

For the inactivation of CSF viruses likely to be present in trophies, one of the following procedures should be used:

1. boiling in water for an appropriate time so as to ensure that any matter other than bone, tusks or teeth is removed;

2. gamma irradiation at a dose of at least 20 kiloGray at room temperature (20°C or higher);

3. soaking, with agitation, in a 4% (w/v) solution of washing soda (sodium carbonate - $\text{Na}_2\text{CO}_3$) maintained at pH 11.5 or above for at least 48 hours;

4. soaking, with agitation, in a formic acid solution (100 kg salt [NaCl] and 12 kg formic acid per 1,000 litres water) maintained at below pH 3.0 for at least 48 hours; wetting and dressing agents may be added;

5. in the case of raw hides, salting for at least 28 days with sea salt containing 2% washing soda (sodium carbonate - $\text{Na}_2\text{CO}_3$).
Annex XXVIII (contd)

Article 15.3.2620.

Surveillance: introduction

Articles 15.3.2620. to 15.3.3425. define the principles and provide a guide on the surveillance for CSF, complementary to Chapter 1.4., applicable to Members seeking to determine their CSF status. This may be for the entire country or a zone. Guidance for Members seeking free status following an outbreak and for the maintenance of CSF status is also provided.

The impact and epidemiology of CSF differ widely in different regions of the world, and it is, therefore, impossible to provide specific recommendations for all situations. The surveillance strategies employed for demonstrating freedom from CSF at an acceptable level of confidence will need to be adapted to the local situation. For example, the approach must be tailored in order to prove freedom from CSF for a country or zone where wild pigs provide a potential reservoir of infection, or where CSF is present in adjacent countries. The method must examine the epidemiology of CSF in the region concerned and adapt to the specific risk factors encountered. This should include provision of scientifically based supporting data. There is, therefore, latitude available to Members to provide a well-reasoned argument to prove that absence of classical swine fever virus (CSFV) infection is assured at an acceptable level of confidence.

Surveillance for CSF should be in the form of a continuing programme designed to establish that a population in a country, zone or compartment is free from CSFV infection or to detect the introduction of CSFV into a population already recognized as free. Consideration should be given to the specific characteristics of CSF epidemiology which include: the role of swill feeding and the impact of different production systems on disease spread, the role of semen in transmission of the virus, the lack of pathognomonic gross lesions and clinical signs, the frequency of clinically inapparent infections, the occurrence of persistent and chronic infections, and the genotypic, antigenic, and virulence variability exhibited by different strains of CSFV. Serological cross-reactivity with other pestiviruses has to be taken into consideration when interpreting data from serological surveys. A common route by which ruminant pestiviruses can infect pigs is the use of vaccines contaminated with bovine viral diarrhoea virus (BVDV).

For the purposes of this Chapter, virus infection means presence of CSFV as demonstrated directly by virus isolation, the detection of virus antigen or virus nucleic acid, or indirectly by seroconversion which is not the result of vaccination.

Article 15.3.2721.

Surveillance: general conditions and methods

1. A surveillance system in accordance with Chapter 1.4. should be under the responsibility of the Veterinary Authority. A procedure should be in place for the rapid collection and transport of samples to an accredited laboratory as described in the Terrestrial Manual.

2. The CSF surveillance programme should:
   a) include an early warning system throughout the production, marketing and processing chain for reporting suspicious cases. Farmers and workers, who have day-to-day contact with livestock, as well as diagnosticians, should report promptly any suspicion of CSF to the Veterinary Authority. They should be supported directly or indirectly (e.g. through private veterinarians or veterinary para-professionals) by government information programmes and the Veterinary Authority. Since many strains of CSFV do not induce pathognomonic gross lesions or clinical signs, cases in which CSF cannot be ruled out should be immediately investigated employing clinical, pathological, and laboratory diagnosis. This requires that sampling kits and other equipment are available to those responsible for surveillance. Personnel responsible for surveillance should be able to call for assistance from a team with expertise in CSF diagnosis, epidemiological evaluation, and control;
b) implement, when relevant, regular and frequent clinical inspections and serological testing of high-risk groups of animals (for example, where swill feeding is practised), or those adjacent to a CSF infected country or zone (for example, bordering areas where infected wild pigs are present).

An effective surveillance system will periodically identify suspicious cases that require follow-up and investigation to confirm or exclude that the cause of the condition is CSFV. The rate at which such suspicious cases are likely to occur will differ between epidemiological situations and cannot, therefore, be reliably predicted. Recognitions for freedom from CSFV infection should, as a consequence, provide details of the occurrence of suspicious cases and how they were investigated and dealt with. This should include the results of laboratory testing and the control measures to which the animals concerned were subjected during the investigation (quarantine, movement standstill orders, etc.).

Article 15.3.2

Surveillance strategies

1. Introduction

There are two basic strategies that can be employed for CSF surveillance depending on the purpose of the Member for seeking recognition of freedom from CSF. In countries historically free of CSF, surveillance programmes should be designed to detect the introduction of CSFV into domestic or wild swine. The optimal strategy to meet this objective is most often targeted surveillance.

The population covered by surveillance aimed at detecting disease and infection should include domestic and wild pig populations within the country or zone to be recognised as free from CSFV infection. Such surveillance may involve opportunistic testing of samples submitted for other purposes, but a more efficient and effective strategy is one which includes targeted surveillance.

Surveillance is targeted to the pig population which presents the highest risk of infection (for example, swill fed farms, pigs reared outdoors or farms in proximity to infected wild pigs). Each Member will need to identify its individual risk factors. These may include: temporal and spatial distribution of past outbreaks, pig movements and demographics, etc.

For reasons of cost, the longevity of antibody levels, as well as the existence of clinically inapparent infections and difficulties associated with differential diagnosis of other diseases, serology is often the most effective and efficient surveillance methodology. In some circumstances, which will be discussed later, clinical and virological surveillance may also have value.

The Member should justify the surveillance strategy chosen as adequate to detect the presence of CSFV infection in accordance with Chapter 1.4. and the epidemiological situation. Cumulative survey results in combination with the results of passive surveillance, over time, will increase the level of confidence in the surveillance strategy. If a Member wishes to apply for recognition by other Members of a specific zone within the country as being free from CSFV infection, the design of the surveillance strategy and the basis for any sampling process would need to be aimed at the population within the zone.

For random surveys, the design of the sampling strategy will need to incorporate epidemiologically appropriate design prevalence. The sample size selected for testing will need to be large enough to detect infection if it were to occur at a predetermined minimum rate. The sample size and expected disease prevalence determine the level of confidence in the results of the survey. The Member must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Chapter 1.4. Selection of the design prevalence in particular clearly needs to be based on the prevailing or historical epidemiological situation.
Irrespective of the survey design selected, the sensitivity and specificity of the diagnostic tests employed are factors in the design, sample size determination and interpretation of the results obtained. Ideally, the sensitivity and specificity of the tests used should be validated for the vaccination/infection history and production class of animals in the target population.

Irrespective of the testing system employed, the surveillance system design should anticipate the occurrence of false positive reactions. This is especially true of the serological diagnosis of CSF because of the recognized cross-reactivity with ruminant pestiviruses. There needs to be an effective procedure for following up positives to ultimately determine with a high level of confidence, whether or not they are indicative of CSFV infection. This should involve confirmatory and differential tests for pestiviruses, as well as further investigations concerning the original sampling unit as well as animals which may be epidemiologically linked.

2. Clinical and virological surveillance

Beyond their role in targeted surveillance, clinical and virological surveillance for CSF has two aims: a) to shorten the period between introduction of CSF virus into a disease free country or zone and its detection, and b) to confirm that no unnoticed outbreaks have occurred.

In the past, clinical identification of cases was the cornerstone of early detection of CSF. However, emergence of low virulence strains of CSF, as well as new diseases - such as post-weaning multisystemic wasting syndrome and porcine dermatitis and nephropathy syndrome - have made such reliance less effective, and, in countries where such diseases are common, can add significant risk of masking the presence of CSF.

The spectrum of disease signs and gross pathology seen in CSF infections, along with the plethora of other agents that can mimic CSF, renders the value of clinical examination alone somewhat inefficient as a surveillance tool. These factors, along with the compounding effects of concurrent infections and diseases caused by ruminant pestiviruses, dictate the need for laboratory testing in order to clarify the status of CSF suspects detected by clinical monitoring.

Nevertheless, clinical presentation should not be ignored as a tool for early detection; in particular, any cases where clinical signs or lesions consistent with CSF are accompanied by high morbidity and/or mortality should be investigated without delay. In CSFV infections involving low virulence strains, high mortality may only be seen in young animals. Otherwise close physical examination of susceptible animals is useful as a selection criteria for CSF surveillance, particularly in diagnostic laboratories or slaughter establishments or when applied to high risk populations such as swill feeding operations.

The difficulties in detecting chronic disease manifested by non-specific clinical signs and delayed seroconversion and seronegativity, in persistently infected piglets, both of which may be clinically normal, makes virological investigation essential. As part of a herd investigation, such animals are likely to be in a minority and would not confound a diagnosis based on serology. Individually or as part of recently mixed batches, such animals may, however, escape detection by this method. A holistic approach to investigation, taking note of herd history, pig, personnel and vehicle movements and disease status in neighbouring zones or countries, can also assist in targeting surveillance in order to increase efficiency and enhance the likelihood of early detection.
Annex XXVIII (contd)

The labour-intensive nature of clinical, pathological and virological investigations, along with the smaller ‘window of opportunity’ inherent in virus, rather than antibody detection, has, in the past, resulted in greater emphasis being placed on mass serological screening as the best method for surveillance. However, surveillance based on clinical and pathological inspection and virological testing should not be underrated. If targeted at high risk groups in particular, it provides an opportunity for early detection that can considerably reduce the subsequent spread of disease. Herds predominated by adult animals, such as nucleus herds and artificial insemination studs, are particularly useful groups to monitor, since infection by low virulence viruses in such groups may be clinically inapparent, yet the degree of spread may be high.

Clinical and virological monitoring may also provide a high level of confidence of rapid detection of disease if a sufficiently large number of clinically susceptible animals is examined. In particular, molecular detection methods are increasingly able to offer the possibility of such large-scale screening for the presence of virus, at reasonable cost.

Wild pigs and, in particular, those with a wholly free-living existence, rarely present the opportunity for clinical observation, but should form part of any surveillance scheme and should, ideally, be monitored for virus as well as antibody.

Vaccine design and diagnostic methodologies, and in particular methods of virus detection, are increasingly reliant on up-to-date knowledge of the molecular, antigenic and other biological characteristics of viruses currently circulating and causing disease. Furthermore, epidemiological understanding of the pathways of spread of CSFV can be greatly enhanced by molecular analyses of viruses in endemic areas and those involved in outbreaks in disease free areas. It is therefore essential that CSFV isolates are sent regularly to the regional OIE Reference Laboratory for genetic and antigenic characterisation.

3. Serological surveillance

Serological surveillance aims at detecting antibodies against CSFV. Positive CSFV antibody test results can have five possible causes:

a) natural infection with CSFV;

b) legal or illegal vaccination against CSF;

c) maternal antibodies derived from an immune sow (maternal antibodies) are usually found only up to 4.5 months of age, but, in some individuals, maternal antibodies can be detected for considerably longer periods;

d) cross-reactions with other pestiviruses;

e) non-specific reactors.

The infection of pigs with other pestiviruses may complicate a surveillance strategy based on serology. Antibodies to bovine viral diarrhoea virus (BVDV) and Border disease virus (BDV) can give positive results in serological tests for CSF, due to common antigens. Such samples will require differential tests to confirm their identity. Although persistently infected immunotolerant pigs are themselves seronegative, they continuously shed virus, so the prevalence of antibodies at the herd level will be high. Chronically infected pigs may have undetectable or fluctuating antibody levels.

It may be possible to use sera collected for other survey purposes for CSF surveillance. However, the principles of survey design described in this Chapter and the requirement for statistical validity should not be compromised.
The discovery of clustering of seropositive reactions should be foreseen. It may reflect any of a series of events, including but not limited to the demographics of the population sampled, vaccinal exposure or the presence of infection by field strains or other pestiviruses. Because clustering may signal field strain infection, the investigation of all instances must be incorporated in the survey design. Clustering of positive animals is always epidemiologically significant and therefore should be investigated.

In countries or zones that are moving towards freedom, serosurveillance can provide valuable information on the disease status and efficacy of any control programme. Targeted serosurveillance of young stock will indicate whether newly circulating virus is present, although the presence of maternal antibody will also need to be considered. If conventional attenuated vaccine is currently being used or has been used in the recent past, serology aimed at detecting the presence of field virus will likewise need to be targeted at unvaccinated animals and after the disappearance of maternal antibody. General usage in such situations may also be used to assess levels of vaccine coverage.

Vaccines also exist which, when used in conjunction with dedicated serological tests, may allow discrimination between vaccinal antibody and that induced by field infection. Such tools, described in the Terrestrial Manual, will need to be fully validated. They do not confer the same degree of protection as that provided by conventional vaccines, particularly with respect to preventing transplacental infections. Furthermore, serosurveillance using such differentiation requires cautious interpretation on a herd basis.

The results of random or targeted serological surveys are important in providing reliable evidence that no CSFV infection is present in a country or zone. It is therefore essential that the survey be thoroughly documented.

Article 15.3.2023

Country or zone historically-free of CSF: additional surveillance procedures

The free status should be reviewed whenever evidence emerges to indicate that changes which may alter the underlying assumption of continuing historical freedom, has occurred. Such changes include but are not limited to:

1. an emergence or an increase in the prevalence of CSF in countries or zones from which live pigs or products are imported;
2. an increase in the volume of imports or a change in their country or zone of origin;
3. an increase in the prevalence of CSF in the domestic or wild pigs of adjacent countries or zones;
4. an increased entry from, or exposure to, infected wild pig populations of adjacent countries or zones.

Article 15.3.2024

Countries, zones or compartments declaring freedom from CSF: additional surveillance procedures

1. Country or zone free of CSF

In addition to the general conditions described in the above-mentioned articles, a Member seeking recognition of CSF freedom for the country or a zone, whether or not vaccination had been practised, should provide evidence for the existence of an effective surveillance programme. The strategy and design of the surveillance programme will depend on the prevailing epidemiological circumstances in and around the country or zone and will be planned and implemented according to the general conditions and methods described in this Chapter, to demonstrate the absence of CSFV infection in domestic and wild pig populations. This requires the support of a national or other laboratory able to undertake identification of CSFV infection through virus detection and serological tests described in the Terrestrial Manual.
Annex XXVIII (contd)

2. **Compartment free of CSF**

The objective of surveillance is to demonstrate the absence of CSFV infection in the compartment. The provisions of Chapter 4.3. should be followed. The effective separation of the two subpopulations should be demonstrated. To this end, a biosecurity plan that includes but is not limited to the following provisions should be implemented:

a) proper containment of domestic pigs;

b) control of movement of vehicles with cleaning and disinfection as appropriate;

c) control of personnel entering into the establishments and awareness of risk of fomite spread;

d) prohibition of introduction to the establishments of wild caught animals and their products;

e) record of animal movements into and out of establishments;

f) information and training programmes for farmers, processors, veterinarians, etc.

The biosecurity plan implemented also requires internal and external monitoring by the Veterinary Authority. This monitoring should include:

a) periodic clinical and serological monitoring of herds in the country or zone, and adjacent wild pig populations following these recommendations;

b) herd registration;

c) official accreditation of biosecurity plans;

d) periodic monitoring and review.

Monitoring the CSF status of wild and domestic pig populations outside the compartment will be of value in assessing the degree of risk they pose to the CSF free compartment. The design of a monitoring system is dependent on several factors such as the size and distribution of the population, the organisation of the Veterinary Services and resources available. The occurrence of CSF in wild and domestic pigs may vary considerably among countries. Surveillance design should be epidemiologically based, and the Member should justify its choice of design prevalence and level of confidence based on Chapter 1.4.

The geographic distribution and approximate size of wild pig populations need to be assessed as a prerequisite for designing a monitoring system. Sources of information may include government wildlife authorities, wildlife conservation organisations, hunter associations and other available sources. The objective of a surveillance programme when the disease is already known to exist should be to determine the geographic distribution and the extent of the infection.

Article 15.3.3125.

**Recovery of free status: additional surveillance procedures**

1. **Countries or zones seeking re-establishment of freedom from CSF following an outbreak**

In addition to the general conditions described in the above-mentioned articles, a Member seeking reestablishment of country or zone freedom from CSF should show evidence of an active surveillance programme to demonstrate absence of CSFV infection.
Annex XXVIII (contd)

Populations under this surveillance programme should include:

a) establishments in the proximity of the outbreak;

b) establishments epidemiologically linked to the outbreak;

c) animals used to re-populate affected establishments and any establishments where contiguous culling is carried out;

d) wild pig populations in the area of the outbreak.

In all circumstances, a Member seeking reestablishment of country or zone freedom from CSF with vaccination or without vaccination should report the results of an active and a passive surveillance programme in which the pig population undergoes regular clinical, pathological, virological, and/or serological examination, planned and implemented according to the general conditions and methods described in these recommendations. The surveillance should be based on a statistically representative sample of the populations at risk.

Article 15.3.26.

Surveillance for CSF in wild pigs

2. Surveillance for CSF in wild pigs

While the same principles apply, surveillance in wild pigs presents challenges beyond those encountered in domestic populations in each of the following areas:

a) determination of the distribution, size and movement patterns associated with the wild pig population;

b) assessment of the possible presence of CSF within the population;

c) determination of the practicability of establishing a zone.

The design of a monitoring system for wild pigs is dependent on several factors such as the organisation of the Veterinary Services and resources available. The geographic distribution and approximate size of wild pig populations need to be assessed as a prerequisite for designing a monitoring system. Sources of information may include wildlife conservation organisations, hunter associations and other available sources. The objective of a surveillance programme is to determine if a given disease is present, and if so, at what prevalence.

Estimates of wild pig populations can be made using advanced methods (e.g., radio tracking, linear transect method, capture/recapture) or traditional methods based on the number of animals that can be hunted to allow for natural restocking (hunting bags).

For implementation of the monitoring programme, it will be necessary to define the limits of the territory over which wild pigs range in order to delineate the epidemiological units within the monitoring programme. It is often difficult to define epidemiological units for wild animals. The most practical approach is based on natural and artificial barriers.
The monitoring programme should also include animals found dead, road kills, animals showing abnormal behaviour or exhibiting gross lesions during dressing.

There may be situations where a more targeted surveillance programme can provide additional assurance. The criteria to define high risk areas for targeted surveillance include:

a) areas with past history of CSF;

b) sub-regions with high [large populations of wild pigs density];

c) border regions with CSF affected countries or zones;

d) interface between wild and domestic pig populations;

e) picnic and camping areas;

f) farms with free-ranging pigs;

g) garbage dumps;

h) other risk areas determined by the Veterinary Authority.
CHAPTER 8.XX.

WEST NILE FEVER

Article 8.XX.1.

General provisions

West Nile fever (WNF) is a zoonotic disease caused by certain strains of the mosquito-borne transmitted West Nile virus (WNV).

For the purpose of this Chapter, the susceptible species are equidae, geese, ducks (under study) and chicken and turkey chicks less than 12 days old (under study) and birds other than poultry.

WNV is maintained in a mosquito–bird–mosquito transmission cycle, whereas humans and equidae are considered dead-end hosts. Most human infections occur by natural transmission from mosquitoes.

In relation to domestic animal trade, geese and ducks pose a risk for the spread of the WNV as some species have been documented to develop a viraemia sufficient to infect mosquitoes.

Surveillance for WNF will should be carried out according to Chapter X.X.

The following criteria define the occurrence of WNF:

1. WNV has been isolated from an animal; or

2. viral antigen or viral ribonucleic acid (RNA) specific to WNV has been identified in samples from one or more animals that show clinical signs consistent with WNF, or that is epidemiologically linked to a confirmed or suspected outbreak of WNF; or

3. antibodies to WNV that are not a consequence of vaccination, have been identified in an animal, that shows clinical signs consistent with WNF, or that is epidemiologically linked to a confirmed or suspected outbreak of WNF.

For the purposes of the Terrestrial Code, the incubation period for WNF shall be 15 days.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 8.XX.1.2.

Trade in commodities

Members should not impose trade restrictions on dead-end hosts such as horses.

When authorising import or transit of the following commodities and any products made from these, Veterinary Authorities should not require any West Nile virus (WNV) related conditions, regardless of the WNF risk status of the animal population of the exporting country or zone.
Annex XXIX (cont’d)

a) hatching eggs;
b) eggs for human consumption;
c) egg products;
d) poultry semen;
e) fresh meat and meat products of poultry;
f) products of poultry origin intended for use in animal feeding, or for agricultural or industrial use;
g) feathers and down from poultry;
h) semen of horses;
i) fresh meat and meat products of horses.

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the WNF status of the ruminant population of the exporting country or zone.

Article 8.XX.21.

West Nile fever (WNF) is a zoonotic disease caused by certain strains of the mosquito-borne West Nile virus (WNV).

For the purpose of this Chapter, the susceptible species are equidae, geese, ducks (under study) and chicken and turkey chicks less than 12 days old and birds other than poultry.

Although most avian species are susceptible to infection, the outcome of the infection is highly variable according to the species. Chickens and turkeys, are usually resistant to disease and do not develop viremia sufficient to infect mosquitoes, with the exception of chicks less than 12 days old.

Birds are responsible for virus dispersal, including reintroduction of WNV from endemic areas into regions that may subsequently experience sporadic outbreaks.

WNV is maintained in a mosquito–bird–mosquito transmission cycle, whereas humans and equidae are considered dead-end hosts. Most human infections occur by natural transmission from mosquitoes.

Many animal species are known to be susceptible to WNV infection and outbreaks of a fatal neurological disease have been reported in humans, equidae, geese and wild birds.

In relation to domestic animal trade, geese and ducks pose a risk for the spread of the WNV as some species have been documented to develop a viremia sufficient to infect mosquitoes.

WNV has been reported to date in a wide geographical range that includes portions of Europe, Asia, Africa, Australia and the Americas. Although competent vectors and susceptible bird species are nearly ubiquitous, WNV circulation in sylvatic cycles may spill over occasionally in domestic population.
Surveillance for WNF will be carried out according to Chapter X.X.

The following criteria defines the occurrence of WNF case:

1. WNV has been isolated and identified as such from an animal, including human; or

2. Viral antigen or viral ribonucleic acid (RNA) specific to WNV has been identified in samples from one or more animals including human that show clinical signs consistent with WNF, or that is epidemiologically linked to a confirmed or suspected outbreak of WNF; or

3. Antibodies to WNV that are not a consequence of vaccination, have been identified in an animal, that including human show clinical signs consistent with WNF, or that is epidemiologically linked to a confirmed or suspected outbreak of WNF.

For the purposes of the Terrestrial Code, the incubation period for WNF shall be 15 days.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 2.2.XX.2.

WNF infected country, or zone or compartment

A WNF infected country, or zone or compartment is a country, zone or compartment clearly defined where one in which a case of WNF has been reported during the past 2 years

Article 8.XX.3.

WNF free country, or zone or compartment

1. A country, or zone or compartment may be considered free from WNF when WNF is notifiable in the whole country and either:

   a) no clinical occurrence of indigenous WNF case, where infection occurred within the territory of the Member, have been recorded for the past 2 years; or

   b) a surveillance programme in accordance with Chapter X.X. has demonstrated no evidence of WNFV in the country or zone or compartment during the past 2 years; or

   c) a surveillance programme has demonstrated no evidence of Culex mosquitoes likely to be competent WNV vectors in the country, or zone or compartment.

2. A WNF free country, or zone or compartment will not lose its free status through the importation from WNF infected countries or infected zones or compartment of:

   a) seropositive animals;

   b) semen, embryo or ova;

   c) animals vaccinated in accordance with the Terrestrial Manual at least 30 days prior to dispatch, and that the animals are identified in the accompanying certification as having been vaccinated; or

   d) animals not vaccinated if a surveillance programme in accordance with Chapter X.X. has been in place in the source population for a period of 30 days immediately prior to dispatch, and no evidence of WNV transmission has been detected.
Annex XXIX (contd)

Article 8.XX.4.

WNF seasonally free country or zone

1. A WNF seasonally free country or zone is a country or a zone for one in which for part of a year, surveillance demonstrates no evidence either of WNV transmission or presence of adult *Culex* mosquitoes likely to be competent WNV vectors.

2. For the application of Article 8.XX.7., the seasonally free period is taken to commence 21 days following the last evidence of WNV transmission (as demonstrated by the surveillance programme), or the cessation of activity of adult *Culex* mosquitoes likely to be competent WNV vectors.

3. For the application of Article 8.XX.7., the seasonally free period is taken to conclude either:
   a) at least 21 days before the earliest date that historical data show WNV transmission cycle has recommenced; or
   b) immediately if current climatic data or data from a surveillance programme indicate an earlier resurgence of activity of adult *Culex* mosquitoes likely to be competent WNV vectors.

4. A WNF seasonally free country or zone will not lose its free status through the importation of animals or semen or embryo and ova from infected countries or zones if:
   a) seropositive animals;
   b) semen, embryo or ova;
   c) animals vaccinated in accordance with the Terrestrial Manual at least 30 days prior to dispatch, and are identified in the accompanying certification as having been vaccinated; or
   d) animals not vaccinated if a surveillance programme in accordance with Chapter X.X. has been in place in the source population for a period of 30 days immediately prior to dispatch, and no evidence of WNV transmission has been detected.

Article 8.XX.5.

WNF infected country or zone

A WNF infected country or zone is one in which a case of WNF has been reported during the past 2 years.

Article 8.XX.6.5

Recommendations for importation from WNF free countries, or zones, or compartment for susceptible species (other than horses)

Veterinary Administrations should require the presentation of an international veterinary certificate attesting that:

1. the animals were kept in a WNF free country, or zone or compartment since birth or for at least 30 days prior to shipment; or
2. the animals were kept in a WNF free country, or zone or compartment, for at least 7–15 days, were subjected, with negative results, to an agent identification test according to the Terrestrial Manual, with negative results, carried out on a sample collected at least 3 days after the commencement of the residence period and remained in the WNF free country, or zone or compartment, until shipment; or

3. the animals:
   a) were vaccinated in accordance with the Terrestrial Manual 30 days before introduction into the free country, or zone or compartment, and
   b) were identified as having been vaccinated; and
   c) were kept in a WNF free country or zone for at least 7–15 days; and
   d) remained in the WNF free country or zone until shipment;

AND

4. if the animals were exported from a WNF free zone, either:
   a) did not transit through an infected country or zone during transportation to the place of shipment; or
   b) were protected from attack from WNV mosquito vectors at all times when transiting through an infected country or zone; or
   c) had been vaccinated in accordance with point 3 above.

Article 8.XX.7.6.

Recommendations for importation from WNF seasonally free countries or zones

for susceptible species (other than horses)

Veterinary Administrations should require the presentation of an international veterinary certificate attesting that the animals:

1. were kept during the seasonally free period in a WNF seasonally free country or zone since birth or for at least 30 days prior to shipment; or

2. were kept during the WNF seasonally free period in a WNF seasonally free country or zone for at least 7–15 days prior to shipment, and were subjected during the residence period in the country or zone to an agent identification test according to the Terrestrial Manual, with negative results, carried out on a sample collected at least 3 days after the commencement of the residence period and remained in the WNF seasonally free country, or zone until shipment; or

3. were kept during the seasonally free period in a WNF seasonally free country or zone for at least 15 days, and were vaccinated in accordance with the Terrestrial Manual 30 days before introduction into the free country or zone against WNF, were identified as having been vaccinated and remained in the WNF seasonally free country or zone until shipment;
AND

4. if the animals were exported from a WNF seasonally free country or zone, either:
   a) did not transit through an infected country or infected zone during transportation to the place of shipment; or
   b) were protected from attack from WNV mosquito vectors at all times when transiting through an infected country or infected zone; or
   c) were vaccinated in accordance with point 3 above.

Article 8.XX.8.7.

Recommendations for importation from WNF infected countries or infected zones

for susceptible species (other than horses)

Veterinary Administrations should require the presentation of an international veterinary certificate attesting that the animals:

1. were protected from attack from WNV mosquito vectors for at least 30 days prior to shipment; or
2. were subjected to a serological test according to the Terrestrial Manual to detect WNV neutralizing antibodies with positive results; or
3. were protected from attack from WNV mosquito vectors for at least 15 days prior to shipment, and were subjected during that period to an agent identification test according to the Terrestrial Manual, with negative results, carried out on a sample collected at least 3 days after being introduced in the mosquito free zone; or
4. were vaccinated in accordance with the Terrestrial Manual at least 30 days before shipment, against WNV, and were identified in the accompanying certification as having been vaccinated; or
5. are not vaccinated and a surveillance programme in accordance with Chapter X.X. has been in place in the source population for a period of 30 days immediately prior to shipment, and no evidence of WNV transmission has been detected;

AND

6. were protected from attack from WNV mosquito vectors during transportation to the place of shipment; or
7. were vaccinated 30 days before shipment or had antibodies against WNV.

Article 8.XX.9.8.

Recommendations for the importation of birds

Veterinary Administrations should require the presentation of an international veterinary certificate attesting that:

1. the birds showed no clinical sign of WNF on the day of shipment; and

Article 8.XX.9.9.
2. the birds were kept in a quarantine station in a mosquito-free environment for 30 days prior to shipment and a statistically valid sample was subjected, with negative results, to an agent identification test according to the Terrestrial Manual at least 3 days after the commencement of the residence period.


Protecting animals from WNV mosquito vectors attacks

When transporting animals through WNF infected countries or infected zones, Veterinary Administrations Authorities should require strategies to protect susceptible animals from attack from WNV mosquitoes vectors during transport, taking into account the local ecology of the vectors mosquitoes.

Potential risk management strategies include:

1. treating animals with chemical insect repellents prior to and during transportation;

2. ensuring vehicles do not stop en route unless the animals are held behind insect proof netting;

3. surveillance for vectors at common stopping and offloading points to gain information on seasonal variations;

4. integrated pest management practices at holding, common stopping and offloading points;

5. using historical, ongoing and/ or WNF modelling information to identify low risk ports and transport routes.

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Introduction

Animal feed is a critical component of the food-chain that has a direct impact on animal health and welfare and also on food safety and public health.

Historically, the OIE primarily addressed animal feed as an important pathway for the entry and spread of contagious epidemic diseases, such as foot and mouth disease, swine vesicular disease and avian influenza. In recent years, the role of feed as a vector for disease agents, including zoonotic organisms, was a focus of standards development in regards to bovine spongiform encephalopathy. Animal feed and feed ingredients are widely traded internationally and trade disruptions have the potential to impact economies in both developed and developing countries. Since 2002 the OIE has expanded its zoonotic disease mandate to encompass animal production food safety, working in collaboration with the Codex Alimentarius Commission (CAC) and other international organisations. In 2006 the International Committee resolved that the OIE should develop guidance on foodborne zoonoses and animal feeding, complementing relevant CAC texts.

Article 2

Purpose Objective and scope

The purpose of this OIE guideline Chapter is to provide guidance on animal feeding in relation to animal health and to complement the guidance provided by the Codex Code of Practice on Good Animal Feeding (CAC/RCP 54-2004) which deals primarily with food safety, and related other Codex texts covering animal feeding, e.g. Code of Practice for Source Directed Measures to Reduce Contamination of Food with Chemicals (CAC/RCP 49-2001).

This guideline This Chapter aims at ensuring the control of animal and public health hazards through adherence to recommended practices during the production (procurement, handling, storage, processing and distribution) and use of both commercial and on-farm produced animal feed and feed ingredients for food-producing terrestrial animals.

Scope

This guideline Chapter applies to the production and use of all products destined for animal feed and feed ingredients at all levels whether produced commercially or on farm. It also includes grazing or free-range feeding, forage crop production and water for drinking. Swill feeding is a particular aspect of on-farm practice that is specifically addressed because of its recognised role in disease transmission.

These Guidelines This Chapter deals with food or feed for terrestrial food-producing animals other than aquatic animals (i.e. livestock and poultry except bees).
Article 3

Definitions

Hazard

Hazard means a biological, chemical or physical agent in, or a condition of, feed or a feed ingredient with the potential to cause an adverse effect on animal or public health.

Feed

Feed means any material (single or multiple), whether processed, semi-processed or raw, which is intended to be fed directly to terrestrial food-producing animals (except bees).

Feed additives

Feed additives means any intentionally added ingredient not normally consumed as feed by itself, whether or not it has nutritional value, which affects the characteristics of feed, or health of the animal or and the characteristics of products of the animal. Microorganisms, enzymes, acidity pH regulators, trace elements, vitamins and other products fall within the scope of this definition depending on the purpose of use and method of administration. This excludes veterinary drugs.

Medicated feed

Medicated feed means any feed which contains a veterinary drug administered to food producing animals, for therapeutic or prophylactic purposes or for modification of physiological functions.

Feed ingredient

Feed ingredient means a component part or constituent of any combination or mixture making up a feed, whether or not it has a nutritional value in the animal’s diet, including feed additives. Ingredients are of plant, (including aquatic plants) or animal terrestrial or aquatic or aquatic animal origin, or other organic or inorganic substances.

Undesirable substance

Undesirable substance means a contaminant or other substance material which is present in and/or on feed and feed ingredients and which constitute a risk whose presence is potentially harmful to animal or public health and/or is restricted under current regulations.

Commercial feed

Commercial feed means all materials that are sold and distributed as feed, or to be mixed with feed, for animals except unmixed seed, whole, processed, or unprocessed; straw, stover, silage, cobs, husks, and hulls; or individual chemical compounds not mixed with other ingredients.

Cross Contamination

Cross Contamination means contamination the presence of a material or product with another material or product containing a component that in a feed or feed ingredient additive and whose presence in that feed or feed ingredient additive is potentially harmful for animal or public health or is restricted under the regulatory framework current regulations.
Article 4

General principles

1. Roles and responsibilities

The Competent Authority has the legal power to set and enforce regulatory animal feeding requirements, and has final responsibility for verifying that these requirements are met. The Competent Authority may establish regulatory requirements for relevant parties to provide it with information and assistance. Refer to Chapters 3.1. and 3.2. of the OIE Terrestrial Code.

Those involved in the production and use of animal feed and feed ingredients have the responsibility to ensure that these products meet regulatory requirements. Appropriate contingency plans should be in place to enable tracing and recall of non-compliant products. All personnel involved in the manufacture, storage and handling of feed and feed ingredients should be adequately trained and aware of their role and responsibility in preventing the introduction or spread of animal health and public health hazards. Appropriate contingency plans should be developed. Equipment manufacturing equipment, storage and transport facilities should be adequate and maintained in good working order and in a sanitary condition.

It is a particular responsibility of Veterinary Services to set and enforce the regulatory requirements pertaining to the use of veterinary drugs, animal disease control and the food safety aspects that relate to the management of live animals on farm.

Those providing specialist services to producers and to the feed industry (e.g. private veterinarians, nutritionists and laboratories) may be required to meet specific regulatory requirements pertaining to the services they provide (e.g. disease reporting, quality standards, transparency).

2. Regulatory safety standards

All feed and feed ingredients should meet regulatory safety standards. In defining limits and tolerances for hazards, scientific evidence, including the sensitivity of analytical methods and on the characterisation of risks, should be taken into account.

3. Risk analysis (risk assessment, risk management and risk communication)

Internationally accepted principles and practices on risk analysis (Section 2 of the OIE Terrestrial Code; and relevant Codex texts) should be used in developing and applying the regulatory framework.

Application of a generic framework should provide a systematic and consistent process for managing all biosecurity risks, while recognising the different risk assessment methodologies used in animal and public health.

4. Good practices

Where national guidelines exist, good agricultural practices and good manufacturing practices (including good hygienic practices) should be followed. Countries without such guidelines are encouraged to develop them.

Where appropriate, Hazard Analysis and Critical Control Point\(^1\) (HACCP) principles should be followed to control hazards that may occur in the manufacture, distribution and feeding of feed and feed additives and feed ingredients.

\(^1\)Hazard Analysis and Critical Control Point, as defined in the Annex to the Recommended International Code of Practice on General Principles of Food Hygiene (CAC/RCP 1-1969).
5. Geographic and environmental considerations

Land and facilities used for production of animal feed and feed ingredients and water sources should not be located in close proximity to epidemiological links between potential sources of hazards for animal health or food safety should be considered when assessing water sources, land or facilities for suitability for the production of animal feed and feed ingredients. Animal health considerations include factors such as disease status, location of quarantined premises and existence of zones/compartments of specified health status. Food safety considerations include factors such as industrial operations that generate pollutants and waste treatment plants.

6. Zoning and compartmentalisation

Feed is an important component of biosecurity and needs to be considered when defining a compartment or zone in accordance with Chapter 4.3. of the OIE Terrestrial Code.

7. Sampling and analysis

Sampling and analytical protocols should be based on scientifically recognised principles and procedures.

8. Labelling

Labelling on how the feed or feed ingredients should be handled, stored and used should be clear and informative as to how the feed and feed ingredients should be handled, stored and used unambiguous, legible and conspicuously placed on the package if sold in package bagged form and on the waybill and other sales documents if sold in bulk, un-packaged bagged form, and should comply with regulatory requirements. See Codex Code of Practice on Good Animal Feeding (CAC/RCP 54-2004), including listing of ingredients and instructions on the handling, storing and use.

9. Design and management of inspection programmes

In meeting animal and public health objectives prescribed in national legislation or required by importing countries, Competent Authorities contribute through the direct performance of some tasks inspection or through the auditing of animal and public health activities conducted by other agencies or the private sector.

Feed and feed ingredients business operators and other relevant parts of industry should practice self-regulation to secure compliance with required standards for procurement, handling, storage, processing, distribution and use. Operators have the primary responsibility for implementing systems for process control. Where such systems are applied. The Competent Authority should verify that they process control systems and safety standards achieve all regulatory requirements.
10. **Assurance and certification**

Feed business operators are responsible for demonstrating the safety of the establishments under their control. Competent Authorities are responsible for providing assurances domestically and to trading partners that regulatory requirements, safety standards have been met. For international trade in animal product based feeds, Veterinary Services are required to provide international veterinary certificates.

11. **Hazards associated with animal feed**

a) **Biological hazards**

Biological hazards that may occur in feed and feed ingredients include agents such as bacteria, viruses, prions, fungi and parasites.

b) **Chemical hazards**

Chemical hazards that may occur in feed and feed ingredients include naturally occurring chemicals (such as mycotoxins and gossypol), industrial and environmental contaminants (such as dioxins and PCBs), residues of veterinary drugs and pesticides and also radionuclides.

c) **Physical hazards**

Physical hazards that may occur in feed and feed ingredients include foreign objects (such as pieces of glass, metal, plastic or wood).

12. **Cross-contamination**

It is important to avoid cross-contamination during the manufacture, storage, distribution (including transport) and use of feed and feed ingredients and relevant provisions should be included in current regulations, the regulatory framework. Scientific evidence, including the sensitivity of analytical methods and on the characterisation of risks, should be drawn upon in developing this framework.

Procedures, such as flushing, sequencing and physical clean-out, should be used to avoid cross-contamination between batches of feed or feed ingredients.

13. **Antimicrobial resistance**

Concerning the use of antimicrobials in animal feed refer to Chapters 6.5. to 6.8. of the OIE Terrestrial Code.

14. **Management of information**

The Competent Authority should establish clear requirements for the provision of information by the private sector as this relates to regulatory requirements.
Records should be maintained in a readily accessible form regarding the production, distribution and use of feed and feed ingredients. These records are required to facilitate the prompt trace-back of feed and feed ingredients to the immediate previous source, and trace-forward to the next subsequent recipients, to address identified animal health or public health concerns (see Section 4.3. of CAC/RCP 54-2004).

Animal identification and animal traceability are tools for addressing animal health (including zoonoses), and food safety risks arising from animal feed (see Chapters 4.1. and 4.2. of the OIE Terrestrial Code; Section 4.3. of CAC/RCP 54-2004).
CHAPTER 8.12.

RIFT VALLEY FEVER

Article 8.12.1.

General provisions

For the purposes of the Terrestrial Code, the infective period for Rift Valley fever (RVF) shall be 30 days.

For the purposes of this Chapter, ruminants include camels.

Standards for diagnostic tests are described in the Terrestrial Manual.

The historic distribution of RVF is the sub-Saharan African continent, Madagascar and the Arabian Peninsula.

Countries or zones within the historic distribution of RVF or adjacent to those that are historically infected should be subjected to surveillance.

Epidemics of RVF may occur in infected areas after flooding. They are separated by inter-epidemic periods that may last for several decades in arid areas and, during these periods, the prevalence of infection in humans, animals and mosquitoes can be difficult to detect.

In the absence of clinical disease, the RVF status of a country or zone within the historically infected regions of the world should be determined by a surveillance programme (carried out in accordance with Chapter 1.4.) focusing on mosquitoes and serology of susceptible mammals. The programme should concentrate on parts of the country or zone at high risk because of historical, geographic and climatic factors, ruminant and mosquito population distribution, and proximity to areas where epidemics have recently occurred.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 8.12.1.bis

Trade in commodities

Commodities other than those listed below are not considered to have the potential to spread RVF when they are the subject of international trade.

When authorising import or transit of the following commodities, Veterinary Authorities should comply with recommendations of this Chapter as relevant to the RVF status of the exporting country or zone:

1. live ruminants;
2. meat and meat products of domestic and wild ruminants;
3. milk and milk products (under study);
Annex XXXI (contd)

When authorising import or transit of the following commodities and any products made from them, Veterinary Authorities should not require any RVF related conditions, regardless of the RVF status of the ruminant population of the exporting country or zone:

1. hides and skins;
2. wool and fiber.

When authorising import or transit of other commodities listed in this Chapter, Veterinary Authorities should require the conditions prescribed in this Chapter relevant to the RVF status of the ruminant population of the exporting country or zone.

Article 8.12.2.

RVF infection free country or zone

A country or a zone may be considered free from RVF infection when the disease is notifiable in animals throughout the country and either:

1. the country or zone lies outside the historically infected regions, and not adjacent to historically infected;
2. a surveillance programme as described in Article 8.12.1. has demonstrated no evidence of RVF infection in humans, animals or mosquitoes in the country or zone during the past 4 years following a RVF epidemic.

The provisions of the last paragraph of Article 8.12.1. may need to be complied with on a continuous basis in order to maintain freedom from infection, depending on the geographical location of the country or zone.

A RVF infection free country or zone in which surveillance and monitoring has found no evidence that RVF infection is present will not lose its free status through the importation of permanently marked seropositive animals or those destined for direct slaughter.

Article 8.12.3.

RVF infected country or zone without disease

A RVF disease free country or zone is a country or zone that is not infection free (see Article 8.12.2.) but in which disease has not occurred in humans or animals in the past 6 months provided that climatic changes predisposing to outbreaks of RVF have not occurred during this time.

Article 8.12.4.

RVF infected country or zone with disease

A RVF infected country or zone with disease is one in which clinical disease in humans or animals has occurred within the past 6 months.
Article 8.12.5.

**Trade in commodities**

Commodities other than those listed below are not considered to have the potential to spread RVF when they are the subject of international trade.

Veterinary Authorities of countries shall consider whether there is a risk with regard to RVF infection in accepting importation or transit through their territory from other countries of the following commodities:

1. live ruminants;
2. meat and meat products of domestic and wild ruminants.

Article 8.12.6.

**Recommendations for importation from RVF infection free countries or zones**

for ruminants

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. were kept in a RVF free country or zone since birth or for at least 30 days prior to shipment; and
2. if the animals were exported from a free zone, either:
   a) did not transit through an infected zone during transportation to the place of shipment; or
   b) were protected from mosquito attack at all times when transiting through an infected zone.

Article 8.12.7.

**Recommendations for importation from RVF infection free countries or zones**

for meat and meat products of domestic and wild ruminants

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the products are derived from animals which remained in the RVF infection free country/free zone since birth or for the last 30 days.

Article 8.12.8.

**Recommendations for importation from RVF infected countries/ zones without disease**

for ruminants

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no evidence of RFV on the day of shipment;
Annex XXXI (contd)

2. met one of the following conditions:
   a) were kept in a RVF infected country/zone free of disease since birth or for the last 6 months providing that climatic changes predisposing to outbreaks of RVF have not occurred during this time; or
   OR
   3. b) were vaccinated against RVF at least 21 days prior to shipment with a modified live virus vaccine; or
   OR
   4. c) were held in a mosquito-proof quarantine station for at least 30 days prior to shipment during which the animals showed no clinical signs of RVF and were protected from mosquitoes between quarantine and the place of shipment as well as at the place of shipment;
   AND
   5. did not transit through an infected zone with disease during transportation of the place of shipment.

Article 8.12.9.

Recommendations for importation from RVF infected countries or zones without disease

for meat and meat products of domestic and wild ruminants

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the products are derived from animals which:
   a) remained in the RVF disease free country/zone infected country or zone without disease since birth or for the last 30 days;
   b) were slaughtered in an approved abattoir and were subjected to ante-mortem and post-mortem inspections for RVF with favourable results;

2. the carcasses from which the products were derived were submitted to maturation at a temperature above +2°C for a minimum period of 24 hours following slaughter.

Article 8.12.10.

Recommendations for importation from RVF infected countries or zones with disease

for ruminants

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no evidence of RVF on the day of shipment;

2. were vaccinated against RVF at least 21 days prior to shipment with a modified live virus vaccine;
Annex XXXI (contd)

OR

3. were held in a mosquito-proof quarantine station for at least 30 days prior to shipment during which the animals showed no clinical signs of RVF and were protected from mosquito attack between quarantine and the place of shipment as well as at the place of shipment.

Article 8.12.11.

Recommendations for importation from RVF infected countries or zones with disease

for meat and meat products of domestic and wild ruminants

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the carcasses:

1. are from animals which have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for RVF with favourable results; and

2. have been fully eviscerated and submitted to maturation at a temperature above +2°C for a minimum period of 24 hours following slaughter.

Article 8.12.12.

Recommendations for importation from RVF infected countries or zones with disease

for in vivo derived embryos of ruminants

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the donor animals:

1. showed no evidence of RVF within the period from 28 days prior to 28 days following collection of the embryos;

2. were vaccinated against RVF at least 21 days prior to collection with a modified live virus vaccine;

OR

3. were serologically tested on the day of collection and at least 14 days following collection and showed no significant rise in titre.

Article 8.12.13. (under study)

Recommendations for importation from RVF infected countries/ zones with or without disease

for milk and milk products
Annex XXXI (contd)

Veterinary authorities of importing countries should require the presentation of an international veterinary certificate attesting that the consignment:

1. was subjected to pasteurization; or

2. was subjected to a combination of control measures with equivalent performance as described in the Codex Alimentarius Code of Hygienic Practice for Milk and Milk Products.

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CHAPTER 11.4.

BOVINE CYSTICERCOSIS

Article 11.4.1.

General provisions

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 11.4.2.

Recommendations for the importation of fresh meat of cattle

Veterinary Authorities of importing countries should require the presentation of an international veterinary certificate attesting that the entire consignment of meat:

1. comes from animals which have been subjected to ante-mortem and post-mortem inspections as described in Chapter 6.2, slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for bovine cysticercosis with favourable results;

2. has been recognised as being free from bovine cysticercosis; or

3. in cases of moderate infestation, has been processed using one of the methods provided in the "Recommended International Code of Practice for ante-mortem and post-mortem judgement of slaughter animals and meat", namely, freezing or heat treatment at 60°C (140°F) (FAO/WHO Codex Alimentarius Commission CAC/RCP 34-1985).

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CHAPTER 15.6.

TESCHOVIRUS ENCEPHALOMYELITIS
(previously enterovirus encephalomyelitis, Teschen
disease, Talfan disease)

Article 15.6.1.

General provisions

For the purposes of the Terrestrial Code, the incubation period for Teschovirus encephalomyelitis shall be 40 days.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 11.8.1 bis

Trade in commodities

Commodities other than those listed below are not considered to have the potential to spread Teschovirus encephalomyelitis when they are the subject of international trade.

When authorising import or transit, Veterinary Authorities should comply with recommendations of this Chapter as relevant to the Teschovirus encephalomyelitis status of the exporting country, zone or compartment.

1. domestic and wild pigs;
2. semen of domestic and wild pigs;
3. fresh meat of domestic and wild pigs;
4. meat products of domestic and wild pigs which have not been processed to ensure the destruction of Teschovirus encephalomyelitis virus;
5. products of animal origin (from pigs) intended for use in animal feeding or for agricultural or industrial use;
6. products of animal origin (from pigs) intended for pharmaceutical or surgical use.

Article 15.6.2.

Teschovirus encephalomyelitis free country

A country may be considered free from Teschovirus encephalomyelitis when it has been shown that Teschovirus encephalomyelitis has not been present for at least the past 3 years.

This period shall be 6 months after the slaughter of the last affected animal for countries in which a stamping-out policy is practised with or without vaccination against Teschovirus encephalomyelitis.
Article 15.6.3.

**Teschovirus encephalomyelitis infected zone**

A zone shall be considered as infected with Teschovirus encephalomyelitis until:

1. at least 40 days have elapsed after the confirmation of the last case and the completion of a stamping-out policy and disinfection procedures, or
2. 6 months have elapsed after the clinical recovery or death of the last affected animal if a stamping-out policy was not practised.

**Article 15.6.4.**

**Trade in commodities**

Veterinary Authorities of Teschovirus encephalomyelitis free countries may prohibit importation or transit through their territory, from countries considered infected with Teschovirus encephalomyelitis, of the following commodities:

1. domestic and wild pigs;
2. semen of domestic and wild pigs;
3. fresh meat of domestic and wild pigs;
4. meat products of domestic and wild pigs which have not been processed to ensure the destruction of Teschovirus encephalomyelitis virus;
5. products of animal origin (from pigs) intended for use in animal feeding or for agricultural or industrial use;
6. products of animal origin (from pigs) intended for pharmaceutical or surgical use.

**Article 15.6.5.**

**Recommendations for importation from Teschovirus encephalomyelitis free countries**

for domestic pigs

the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of Teschovirus encephalomyelitis on the day of shipment;
2. were kept in a country free from Teschovirus encephalomyelitis since birth or for at least the past 40 days.
Article 15.6.6.

**Recommendations for importation from Teschovirus encephalomyelitis free countries**

_for wild pigs_

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of Teschovirus encephalomyelitis on the day of shipment;
2. come from a country free from Teschovirus encephalomyelitis;

_if the country of origin has a common border with a country considered infected with Teschovirus encephalomyelitis:

3. were kept in a quarantine station for the 40 days prior to shipment._

Article 15.6.7.

**Recommendations for importation from countries considered infected with Teschovirus encephalomyelitis**

_for domestic pigs_

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of Teschovirus encephalomyelitis on the day of shipment;
2. were kept since birth, or for the past 40 days, in an establishment where no case of Teschovirus encephalomyelitis was officially reported during that period, and that the establishment of origin was not situated in an Teschovirus encephalomyelitis infected zone; or
3. were kept in a quarantine station for the 40 days prior to shipment;
4. have not been vaccinated against Teschovirus encephalomyelitis; or
5. were vaccinated against Teschovirus encephalomyelitis, not less than 30 days and not more than one year prior to shipment (the nature of the vaccine used, whether inactivated or modified live virus, and the virus types and strains included shall also be stated in the certificate).

Article 15.6.8.

**Recommendations for importation from countries considered infected with Teschovirus encephalomyelitis**

_for wild pigs_

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the animals:
1. showed no clinical sign of Teschovirus encephalomyelitis on the day of shipment;
2. were kept in a quarantine station for the 40 days prior to shipment;
3. have not been vaccinated against Teschovirus encephalomyelitis; or
4. were vaccinated against Teschovirus encephalomyelitis, not less than 30 days and not more than one year prior to shipment (the nature of the vaccine used, whether inactivated or modified live virus, and the virus types and strains included shall also be stated in the certificate).

Article 15.6.9.

Recommendations for importation from Teschovirus encephalomyelitis free countries for semen of pigs

Veterinary authorities should require the presentation of an international veterinary certificate attesting that the donor animals:

1. showed no clinical sign of Teschovirus encephalomyelitis on the day of collection of the semen;
2. were kept in a country free from Teschovirus encephalomyelitis for not less than 40 days prior to collection.

Article 15.6.10.

Recommendations for importation from countries considered infected with Teschovirus encephalomyelitis for semen of pigs

Veterinary authorities should require the presentation of an international veterinary certificate attesting that the donor animals:

1. showed no clinical sign of Teschovirus encephalomyelitis on the day of collection of the semen;
2. were kept in the exporting country, for the 40 days prior to collection, in an establishment or artificial insemination centre where no case of Teschovirus encephalomyelitis was officially reported during that period, and that the establishment or artificial insemination centre was not situated in an Teschovirus encephalomyelitis infected zone.

Article 15.6.11.

Recommendations for importation from Teschovirus encephalomyelitis free countries for fresh meat of pigs

Veterinary authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals:

1. which have been kept in a country free from Teschovirus encephalomyelitis since birth or for at least the past 40 days;
2. which have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for Teschovirus encephalomyelitis with favourable results.

Article 15.6.12.

Recommendations for importation from countries considered infected with Teschovirus encephalomyelitis

for fresh meat of pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals:

1. which have not been kept in an Teschovirus encephalomyelitis infected zone;
2. which have been slaughtered in an approved abattoir not situated in an Teschovirus encephalomyelitis infected zone and have been subjected to ante-mortem and post-mortem inspections for Teschovirus encephalomyelitis with favourable results.

Article 15.6.13.

Recommendations for importation from countries considered infected with Teschovirus encephalomyelitis

for meat products of pigs

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that:

1. the entire consignment of meat products comes from animals which have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for Teschovirus encephalomyelitis with favourable results;
2. the meat products have been processed to ensure the destruction of the Teschovirus encephalomyelitis virus;
3. the necessary precautions were taken after processing to avoid contact of the meat with any source of Teschovirus encephalomyelitis virus.

Article 15.6.14.

Recommendations for importation from Teschovirus encephalomyelitis free countries

for products of animal origin (from pigs) intended for use in animal feeding or for agricultural or industrial use

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products come from animals which have been kept in a country free from Teschovirus encephalomyelitis since birth or for at least the past 40 days.
Article 15.6.15.

**Recommendations for importation from countries considered infected with Teschovirus encephalomyelitis**

for meal and flour from blood, meat, defatted bones, hooves and claws (from pigs)

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products have been processed using heat treatment to ensure the destruction of Teschovirus encephalomyelitis virus.

Article 15.6.16.

**Recommendations for importation from countries considered infected with Teschovirus encephalomyelitis**

for bristles

Veterinary Authorities should require the presentation of an international veterinary certificate attesting that these products have been processed to ensure the destruction of Teschovirus encephalomyelitis virus, in premises controlled and approved by the Veterinary Authority of the exporting country.
CHAPTER 1.5.

STATUS FOR OIE LISTED DISEASES: PROCEDURES FOR SELF DECLARATION AND FOR OFFICIAL RECOGNITION BY THE OIE

Article 1.

General principles

Members may wish to make a self declaration as to the freedom of a country, zone or compartment from an OIE listed disease. The Member may inform the OIE of its claimed status and the OIE may publish the claim. Publication does not imply endorsement of the claim. Self declaration does not apply to bovine spongiform encephalopathy (BSE), foot and mouth disease (FMD), rinderpest and contagious bovine pleuropneumonia (CBPP).

Members may request official recognition by the OIE as to:

1. the risk status of a country or zone with regard to BSE;
2. the freedom of a country or zone from FMD, with or without vaccination;
3. the freedom of a country from rinderpest;
4. the freedom of a country or zone from CBPP.

The OIE does not grant official recognition for other diseases.

In all cases, Members should present documentation setting out the compliance of the Veterinary Services of the applicant country or zone with the provisions of Chapters 1.1., 3.1. and 3.2. of the Terrestrial Code and with the provisions of the relevant disease chapters in the Terrestrial Code and the Terrestrial Manual.

When requesting official recognition of disease status, the Member should submit to the OIE Scientific and Technical Department a dossier providing the information requested (as appropriate) in Articles 2 (for BSE), 3 (for FMD), 4 (for rinderpest) or 5 (for CBPP).

The OIE framework for the official recognition and maintenance of disease status is described in Resolution XXII (administrative procedures) and Resolution XXIII (financial obligations) adopted during the 76th General Session in May 2008.

Article 2.

BSE QUESTIONNAIRE

General introduction

Acceptance of this submission is based on the compliance of the Veterinary Service of the applicant country, zone or compartment with the provisions of Chapters 3.1. of the Terrestrial Code and the compliance of BSE diagnostic laboratories with the provisions of Chapter 1.1.3. of the Terrestrial Manual. Documentary evidence should be provided to support this based on Chapter 3.2. of the Terrestrial Code.
Annex XXXII (contd)

The OIE Terrestrial Code Chapter on BSE, Article 11.6.2, prescribes the criteria to determine the BSE risk status of a the cattle population of a country, zone or compartment. This document is the means whereby a claim for negligible risk (Article 11.6.3.) or controlled risk (Article 11.6.4.) can be made to the OIE.

The document comprises the following:
Section 1 - Risk assessment (Article 11.6.2. § 1)
Section 2 - Other requirements of Article 11.6.2. §2-4
• Ongoing awareness program
• Compulsory notification and investigation
• Diagnostic capability

Section 3 - Surveillance (Article 11.6.2. and Articles 11.6.20. to 11.6.22.)

Section 4 - BSE history of the country, zone or compartment (Article 11.6.3. and 11.6.4.)

N.B. Where, during the completion of this questionnaire, the submitting Veterinary Service provides documentation regarding the legislation under which it is mandated, it should provide the content of any legal act described (in one of the three official languages of OIE), as well as the dates of official publication and implementation. Submitting countries are encouraged to follow the format and numbering used in this document.

SECTION 1: RISK ASSESSMENT (11.6.2.§1)

Introduction

The first step in determining the bovine spongiform encephalopathy (BSE) risk status of the cattle population of a country, zone or compartment is to conduct a risk assessment (reviewed annually), based on Section 2. and 3. and Chapter 4.3. of the Terrestrial Code, identifying all potential factors for BSE occurrence and their historic perspective.

Documentation guidelines

This section provides guidance on the data gathering and presentation of information required to support the risk release and exposure assessments in respect of:

Release assessment
1. The potential for the release of the BSE agent through importation of meat-and-bone meal or greaves
2. The potential for the release of the BSE agent through the importation of potentially infected live cattle
3. The potential for the release of the BSE agent through the importation of potentially infected products of bovine origin

Exposure assessment
4. The origin of bovine carcasses, by-products and slaughterhouse waste, the parameters of the rendering processes and the methods of cattle feed production
5. The potential for the exposure of cattle to the BSE agent through consumption of meat-and-bone meal or greaves of bovine origin

In each of the five areas of release and exposure assessment that follow, the contributor is guided in terms of the question, the rationale and the evidence required to support the country, zone or compartment status claim.
Release assessment

1.1. The potential for the release of the BSE agent through importation of meat-and-bone meal or greaves

Question to be answered: Has meat-and-bone meal, greaves, or feedstuffs containing either, been imported within the past 8 years? If so, where from and in what quantities?

Rationale: Knowledge of the origin of meat-and-bone meal, greaves or feedstuffs containing either meat-and-bone meal or greaves, is necessary to assess the risk of release of BSE agent. Meat-and-bone meal and greaves originating in countries of high BSE risk pose a higher release risk than that from low risk countries. Meat-and-bone meal and greaves originating in countries of unknown BSE risk pose an unknown release risk.

This point is irrelevant if the exposure assessment outlined below in Article 11.6.27. indicates that meat-and-bone meal or greaves has not been fed, either deliberately or accidentally, in the past 8 years. Nevertheless, documentation should be provided on the control systems (including relevant legislation) in place to ensure that meat-and-bone meal or greaves has not been fed to cattle.

Evidence required:

1.1.1. Documentation to support claims that meat-and-bone meal, greaves or feedstuffs containing either meat-and-bone meal or greaves have not been imported, OR

1.1.2. Documentation on annual volume, by country of origin, of meat-and-bone meal, greaves or feedstuffs containing them imported during the past 8 years.

1.1.3. Documentation describing the species composition of the imported meat-and-bone meal, greaves or feedstuffs containing them.

1.1.4. Documentation, from the Veterinary Service of the country of production, supporting why the rendering processes used to produce meat-and-bone meal, greaves or feedstuffs containing them would have inactivated, or significantly reduced the titre of BSE agent, should it be present.

1.2. The potential for the release of the BSE agent through the importation of potentially infected live cattle

Question to be answered: Have live cattle been imported within the past 7 years?

Rationale: The release risks are dependent on:

• country, zone or compartment of origin and its BSE status, which will change as more data become available; this may result from the detection of clinical disease, or following active surveillance, or assessment of geographical BSE risk;

• feeding and management of the imported cattle in the country, zone or compartment of origin;

• use to which the commodity has been put as apart from representing risk of developing clinical disease, the slaughter, rendering and recycling in meat-and-bone meal of imported cattle represents a potential route of exposure of indigenous livestock even if meat-and-bone meal and greaves, or feedstuffs containing them, have not been imported;

• dairy versus meat breeds, where there are differences in exposure in the country, zone or compartment of origin because feeding practices result in greater exposure of one category;
Annex XXXII (contd)

- age at slaughter.

Evidence required:

1.2.1. Documentation including tables on the country, zone or compartment of origin of imports. This should identify the country, zone or compartment of origin of the cattle, the length of time they lived in that country, zone or compartment and of any other country in which they have resided during their lifetime.

1.2.2. Documentation including tables describing origin and volume of imports.

1.2.3. Documentation demonstrating that risks are periodically reviewed in light of evolving knowledge on the BSE status of the country, zone or compartment of origin.

1.3. The potential for the release of the BSE agent through the importation of potentially infected products of bovine origin

Question to be answered: What products of bovine origin have been imported within the past 7 years?

Rationale: The release risks are dependent on:

- the origin of the cattle products and whether these products contain tissues known to contain BSE infectivity (Article 11.6.13.);
- country, zone or compartment of origin and its BSE status, which will change as more data become available; this may result from the detection of clinical disease, or following active surveillance, or assessment of geographical BSE risk;
- feeding and management of the cattle in the country, zone or compartment of origin;
- use to which the commodity has been put apart from representing risk of developing clinical disease, the slaughter, rendering and recycling in meat-and-bone meal of imported cattle represents a potential route of exposure of indigenous livestock even if meat-and-bone meal and grays, or feedstuffs containing them, have not been imported;
- dairy versus meat breeds, where there are differences in exposure in the country, zone or compartment of origin because feeding practices result in greater exposure of one category;
- age at slaughter.

Evidence required:

1.3.1. Documentation on the country, zone or compartment of origin of imports. This should identify the country, zone or compartment of origin of cattle from which the products were derived, the length of time they lived in that country, zone or compartment and of any other country in which they have resided during their lifetime.

1.3.2. Documentation describing origin and volume of imports

1.3.3. Documentation demonstrating that risks are periodically reviewed in light of evolving knowledge on the BSE status of the country, zone or compartment of origin.
Exposure assessment

1.4. The origin of bovine carcasses, by-products and slaughterhouse waste, the parameters of the rendering processes and the methods of cattle feed production

Question to be answered: How have bovine carcasses, by-products and slaughterhouse waste been processed over the past 8 years?

Rationale: The overall risk of BSE in the cattle population of a country, zone or compartment is proportional to the level of known or potential exposure to BSE infectivity and the potential for recycling and amplification of the infectivity through livestock feeding practices. For the risk assessment to conclude that the cattle population of a country, zone or compartment is of negligible or controlled BSE risk, it must have demonstrated that appropriate measures have been taken to manage any risks identified. If potentially infected cattle or contaminated materials are rendered, there is a risk that the resulting meat-and-bone meal could retain BSE infectivity. Where meat-and-bone meal is utilized in the production of any cattle feed, the risk of cross-contamination exists.

Evidence required:

1.4.1. Documentation describing the collection and disposal of fallen stock and materials condemned as unfit for human consumption.

1.4.2. Documentation including tables describing the fate of imported cattle, including their age at slaughter or death.

1.4.3. Documentation describing the definition and disposal of specified risk material, if any.

1.4.4. Documentation describing the rendering process and parameters used to produce meat-and-bone meal and greaves.

1.4.5. Documentation describing methods of animal feed production, including details of ingredients used, the extent of use of meat-and-bone meal in any livestock feed, and measures that prevent cross-contamination of cattle feed with ingredients used in monogastric feed.

1.4.6. Documentation describing the end use of imported cattle products and the disposal of waste.

1.4.7. Documentation describing monitoring and enforcement of the above

1.5. The potential for the exposure of cattle to the BSE agent through consumption of meat-and-bone meal or greaves of bovine origin

Question to be answered: Has meat-and-bone meal or greaves of bovine origin been fed to cattle within the past 8 years (Articles 11.6.3. and 11.6.4. in the Terrestrial Code)?

Rationale: If cattle have not been fed products of bovine origin (other than milk or blood) potentially containing meat-and-bone meal or greaves of bovine origin within the past 8 years, meat-and-bone meal and greaves can be dismissed as a risk.

In the case of countries applying for negligible risk status, it will be required to demonstrate that the ruminant feed ban has been effective for at least 8 years following the birth of the youngest case.
Annex XXXII (contd)

Evidence required:

1.5.1. Documentation describing the use of imported meat-and-bone meal and greaves, including the feeding of any animal species.

1.5.2. Documentation describing the use made of meat-and-bone meal and greaves produced from domestic cattle, including the feeding of any animal species.

1.5.3. Documentation on the measures taken to control cross-contamination of cattle feedstuffs with the meat-and-bone meal and greaves including the risk of cross-contamination during production, transport, storage and feeding.

1.5.4a) Documentation, in the form of the following table, on the audit findings in rendering plants and feed mills processing ruminant material or mixed species containing ruminant material, related to the prohibition of the feeding to ruminants of meat-and-bone meal and greaves.

<table>
<thead>
<tr>
<th>Year (information should be provided for each of the 8 years for effectiveness is claimed)</th>
<th>Type of plant (renderer or feed mill)</th>
<th>Number of plants processing ruminant material</th>
<th>Number of plants in (A) inspected</th>
<th>Total number of visual inspections in (B)</th>
<th>Total number of plants in (B) with infractions</th>
<th>Total number of inspected plants in (B) with sampling</th>
<th>Total number of plants in (C) with positive test results</th>
</tr>
</thead>
<tbody>
<tr>
<td>(A)</td>
<td>(B)</td>
<td>(C)</td>
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<td>Year 1</td>
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<td>Feed mill</td>
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<td>Year 2 etc.</td>
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<td>Feed mill</td>
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</table>

1.5.4b) Documentation, in the form of the following table, on the audit findings in rendering plants and feed mills processing non-ruminant material, related to the prohibition of the feeding of meat-and-bone meal and greaves to ruminants.

<table>
<thead>
<tr>
<th>Year (information should be provided for each of the 8 years for effectiveness is claimed)</th>
<th>Type of plant (renderer or feed mill)</th>
<th>Number of plants processing non-ruminant material</th>
<th>Number of plants in (A) inspected</th>
<th>Total number of visual inspections in (B)</th>
<th>Total number of plants in (B) with infractions</th>
<th>Total number of inspected plants in (B) with sampling</th>
<th>Total number of plants in (C) with positive test results</th>
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<tr>
<td>(A)</td>
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<td>Year 1</td>
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<td>Feed mill</td>
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<td>Year 2 etc.</td>
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<td>Feed mill</td>
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</table>
Annex XXXII (contd)

1.5.5a) Documentation, in the form of the following table, on each plant above processing ruminant material or mixed species containing ruminant material with infractions, specifying the type of infraction and the method of resolution.

<table>
<thead>
<tr>
<th>Year (information should be provided for each of the 8 years for effectiveness is claimed)</th>
<th>Type of plant (renderer or feed mill)</th>
<th>Plant ID</th>
<th>Nature of infraction</th>
<th>Method of resolution</th>
<th>Follow up results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1</td>
<td>Renderer</td>
<td>ID 1</td>
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<td>ID 2</td>
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<td>ID 3 etc.</td>
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<td>Feed mill</td>
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<td>ID 2</td>
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<td>ID 3 etc.</td>
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<td>Year 2 etc.</td>
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</tbody>
</table>

1.5.5b) Documentation, in the form of the following table, on each plant above processing non-ruminant material with infractions, specifying the type of infraction and the method of resolution.

<table>
<thead>
<tr>
<th>Year (information should be provided for each of the 8 years for effectiveness is claimed)</th>
<th>Type of plant (renderer or feed mill)</th>
<th>Plant ID</th>
<th>Nature of infraction</th>
<th>Method of resolution</th>
<th>Follow up results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1</td>
<td>Renderer</td>
<td>ID 1</td>
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<td>ID 2</td>
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<td>ID 3 etc.</td>
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<td></td>
<td>Feed mill</td>
<td>ID 1</td>
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<td>ID 2</td>
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<td></td>
<td>ID 3 etc.</td>
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<td>Year 2 etc.</td>
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<td>Feed mill</td>
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</tbody>
</table>

1.5.6. Documentation explaining why, in light of the findings displayed in the preceding four tables, it is considered that there has been no significant exposure of cattle to the BSE agent through consumption of meat-and-bone meal or greaves of bovine origin.

1.5.7. Documentation of husbandry practices (multiple species farms) which could lend themselves to cross-contamination of cattle feed with meat-and-bone meal and greaves destined to other species.
SECTION 2: OTHER REQUIREMENTS (11.6.2. § 2-4)

2.1. Awareness program (Article 11.6.2. § 2)

Questions to be answered:

- Is there an awareness programme?
- What is the target audience?
- What is the curriculum and how long has it been in place?
- Is there a contingency and/or preparedness plan that deals with BSE?

Rationale

An awareness program is essential to ensure detection and reporting of BSE, especially in countries of low prevalence and competing differential diagnoses.

Evidence required

2.1.1. Documentation indicating when the awareness program was instituted and its continuous application and geographical coverage.

2.1.2. Documentation on the number and occupation of persons who have participated in the awareness program (veterinarians, producers, workers at auctions, slaughterhouses, etc.)

2.1.3. Documentation of materials used in the awareness program (the manual, supportive documents, or other teaching materials).

2.1.4. Documentation on the contingency plan.

2.2. Compulsory notification and investigation (Article 11.6.2. § 3)

Questions to be answered:

- What guidance is given to veterinarians, producers, workers at auctions, slaughterhouses, etc.) in terms of the criteria that would initiate the investigation of an animal as a BSE suspect? Have these criteria evolved?
- What were the date and content of the legal act making notification of BSE suspects compulsory?
- What are the measures in place to stimulate notification, such as compensation payments or penalties for not notifying a suspect?

Rationale

The socio-economic implications associated with BSE require that there be incentives and/or obligations to notify and investigate suspect cases.
Evidence required

2.2.1. Documentation on the date of official publication and implementation of compulsory notification. Including a brief description of incentives and penalties.

2.2.2. Documentation on the manual of procedures for investigation of suspect animals and follow-up of positive findings.

2.3. Examination in an approved laboratory of brain or other tissues collected within the framework of the aforementioned surveillance system (Article 11.6.2. § 5)

Questions to be answered:

- Are the diagnostic procedures and methods those described in Chapter 2.4.6. of the Terrestrial Manual?
- Have these diagnostic procedures and methods been applied through the entire surveillance period?

Rationale

The OIE only recognizes for the purpose of this submission samples that have been tested in accordance with the Terrestrial Manual.

Evidence required

2.3.1. Documentation as to the approved laboratories where samples of cattle tissues from the country, zone or compartment are examined for BSE. (If this is located outside the country, information should be provided on the cooperation agreement).

2.3.2. Documentation of the diagnostic procedures and methods used.

2.3.3. Documentation that the diagnostic procedures and methods have been applied through the entire surveillance period.

SECTION 3: BSE SURVEILLANCE AND MONITORING SYSTEM (11.6.2. § 4)

Questions to be answered:

- Does the BSE surveillance programme comply with the guidelines in Articles 11.6.20. to 11.6.22. of the Terrestrial Code?
- What were the results of the investigations?

Rationale

Article 11.6.2.§ 4 and Articles 11.6.20. to 11.6.22. prescribe the number of cattle, by subpopulation, that need to be tested in order to ensure the detection of BSE at or above a minimal threshold prevalence.

Evidence required

3.1. Documentation that the samples collected are representative of the distribution of cattle population in the country, zone or compartment.
Annex XXXII (cont'd)

3.2. Documentation of the methods applied to assess the ages of animals sampled and the proportions for each method (individual identification, dentition, other methods to be specified).

3.3. Documentation of the means and procedures whereby samples were assigned to the cattle subpopulations described in 11.6.21., including the specific provisions applied to ensure that animals described as clinical met the conditions of 11.6.21§1.

3.4. Documentation of the number of animals meeting 11.6.21§1 as compared to the numbers of clinical samples submitted in previous years in accordance to the former provisions in the Code, and explanation of possible differences.

3.5. Documentation, based on the following table, of all clinically suspect cases notified complying with the definition in 11.6.21§1.

<table>
<thead>
<tr>
<th>Laboratory identification number</th>
<th>Age</th>
<th>Clinical signs</th>
<th>Point of detection (farm, market channels, slaughterhouse)</th>
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</thead>
<tbody>
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</tbody>
</table>

3.6. Documentation according to the following table, that the number of target points applicable to the country, zone or compartment and its BSE surveillance requirements (Type A or type B surveillance as a result of the risk assessment of section 1) are met as described in 11.6.21 and 11.6.22.

**SUMMARY TABLE FOR BSE SURVEILLANCE**

<table>
<thead>
<tr>
<th>Year: (complete a separate table for each year of surveillance)</th>
<th>Surveillance subpopulations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Routine slaughter</td>
</tr>
<tr>
<td></td>
<td>Samples</td>
</tr>
<tr>
<td>&gt;1 and &lt;2 years</td>
<td></td>
</tr>
<tr>
<td>=2 and &lt;4 years</td>
<td></td>
</tr>
<tr>
<td>=4 and &lt;7 years</td>
<td></td>
</tr>
<tr>
<td>=7 and &lt;9 years</td>
<td></td>
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<tr>
<td>=9 years</td>
<td></td>
</tr>
<tr>
<td>Subtotals</td>
<td></td>
</tr>
<tr>
<td>Total points</td>
<td></td>
</tr>
</tbody>
</table>

3.7. Indicate the number of adult cattle (over 24 month of age) in the country, zone or compartment.
SECTION 4: BSE HISTORY OF THE COUNTRY, ZONE OR COMPARTMENT (11.6.3. and 11.6.4.)

Questions to be answered

- Has BSE occurred in the country, zone or compartment?
- How has it been dealt with?

Rationale

The categorization of a country, zone or compartment in either negligible or controlled risk is dependent upon, the outcome of the risk assessment described in section 1, compliance with the provisions described in section 2, the results of surveillance described in section 3, and the history of BSE in the country, zone or compartment. This section provides the opportunity to describe the BSE history in the country, zone or compartment.

Evidence required

4.1. Documentation of whether a case of BSE has ever been diagnosed in the country, zone or compartment.

In the case of positive BSE findings:

4.2. Documentation on the origin of each BSE case in respect to the country, zone or compartment. Indicate the birth date and place of birth.

4.3. Indicate the most recent year of birth in relation to all BSE cases

4.4. Documentation that:

the case(s) and all the progeny of female cases, born within 2 years prior to or after clinical onset of the disease, and

all cattle which, during their first year of life, were reared with the BSE cases during their first year of life, and which investigation showed consumed the same potentially contaminated feed during that period, or

if the results of the investigation are inconclusive, all cattle born in the same herd as, and within 12 months of the birth of, the BSE cases,

if alive in the country, zone or compartment, are permanently identified, and their movements controlled, and, when slaughtered or at death, are completely destroyed.
Article 3.

**FMD QUESTIONNAIRE**

**FMD FREE COUNTRY WHERE VACCINATION IS NOT PRACTISED**

Report of Country which applies for recognition of status, under Chapter 8.5. of the Terrestrial Animal Health Code 2008, as a FMD free country not practising vaccination

Please address concisely the following topics. National regulations laws and Veterinary Administration directives may be referred to and annexed as appropriate in one of the OIE official languages.

1. **Introduction**
   1.1. Geographical factors. Provide a general description of the country including physical, geographical and other factors that are relevant to FMD dissemination, countries sharing common borders and other countries that although may not be adjacent share a link for the potential introduction of disease. Provide a map identifying the factors above.
   1.2. Livestock industry. Provide a general description of the livestock industry in the country.

2. **Veterinary system**
   2.1. Legislation. Provide a list and summary of all relevant veterinary legislation in relation to FMD.
   2.2. Veterinary Services. Provide documentation on the compliance of the Veterinary Service of the country with the provisions of Chapters 3.1. and 3.2. of the Terrestrial Code and 1.1.3. of the Terrestrial Manual and describe how the veterinary services supervise and control all FMD related activities. Provide maps and tables wherever possible.
   2.3. Role of farmers, industry and other relevant groups in FMD surveillance and control (include a description of training and awareness programs on FMD).
   2.4. Role of private veterinary profession in FMD surveillance and control.

3. **FMD eradication**
   3.1. History. Provide a description of the FMD history in the country, date of first detection, origin of infection, date of eradication (date of last case), types and subtypes present.
   3.2. Strategy. Describe how FMD was controlled and eradicated (e.g. stamping-out, modified stamping-out, zoning), provide timeframe for eradication.
   3.3. Vaccines and vaccination. Was FMD vaccine ever used? If so, when was the last vaccination carried out? What species were vaccinated?
   3.4. Legislation, organisation and implementation of the FMD eradication campaign. Provide a description of the organizational structure at the different levels. Indicate if detailed operational guidelines exist and give a brief summary.
3.5. Animal identification and movement control. Are susceptible animals identified (individually or at a group level)? Provide a description of the methods of animal identification, herd registration and traceability. How are animal movements controlled in the country? Provide evidence on the effectiveness of animal identification and movement controls. Please provide information on pastoralism, transhumance and related paths of movement.

4. FMD diagnosis

Provide documentary evidence that the provisions in Chapters 1.1.2., 1.1.3., and 2.1.5. of the Terrestrial Manual are applied. In particular, the following points should be addressed:

4.1. Is FMD laboratory diagnosis carried out in the country? If so, provide a list of approved laboratories. If not, provide the name(s) of and the arrangements with the laboratory(ies) samples are sent to, the follow up procedures and the time frame for obtaining results.

4.2. Provide an overview of the FMD approved laboratories, in particular to address the following points:

4.2.1. Procedures for the official accreditation of laboratories. Give details of internal quality management systems, e.g. Good Laboratory Practice, ISO etc. that exist in, or planned for, the laboratory system.

4.2.2. Give details of participation in inter-laboratory validation tests (ring tests).

4.2.3. Is live virus handled?

4.2.4. Biosecurity measures applied.

4.2.5. Details of the type of tests undertaken.

5. FMD surveillance

Provide documentary evidence that surveillance for FMD in the country complies with the provisions of Articles 8.5.40. to 8.5.46. of the Terrestrial Code and Chapter 2.1.5. of the Terrestrial Manual. In particular, the following points should be addressed:

5.1. Clinical suspicion. What are the criteria for raising a suspicion of FMD? What is the procedure to notify (by whom and to whom) and what penalties are involved for failure to report? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for FMDV, species, type of sample, testing method(s) and results (including differential diagnosis).

5.2. Serological surveillance. Are serological surveys conducted? If so, provide detailed information on the survey design (confidence level, sample size, stratification) How frequently are they conducted? Are wildlife susceptible species included in serological surveys? Provide a summary table indicating, for the past two years, the number of samples tested for FMDV, species, type of sample, testing method(s) and results (including differential diagnosis). Provide details on follow up actions taken on all suspicious and positive results. Provide criteria for selection of populations for targeted surveillance and numbers of animals examined and samples tested. Provide details on the methods applied for monitoring the performance of the surveillance system including indicators.
5.3. Livestock demographics and economics. What is the susceptible animal population by species and production systems? How many herds, flocks, etc., of each susceptible species are in the country? How are they distributed (e.g., herd density, etc.)? Provide tables and maps as appropriate.

5.4. Wildlife demographics. What susceptible species are present in the country? Provide estimates of population sizes and geographic distribution. What are the measures in place to prevent contact between domestic and wildlife susceptible species?

5.5. Slaughterhouses and markets. Where are the major livestock marketing or collection centers? What are the patterns of livestock movement within the country? How are the animals transported and handled during these transactions.

6. FMD prevention

6.1. Coordination with neighbouring countries. Are there any relevant factors about the adjacent countries or zones that should be taken into account (e.g., size, distance from adjacent border to affected herds or animals)? Describe coordination, collaboration and information sharing activities with neighboring countries.

6.2. Import control procedures

From what countries or zones does the country authorize the import of susceptible animals or their products? What criteria are applied to approve such countries or zones? What controls are applied on entry of such animals and products, and subsequent internal movement? What import conditions and test procedures are required? Are imported animals of susceptible species required to undergo a quarantine or isolation period? If so, for how long and where? Are import permits and health certificates required? What other procedures are used? Provide summary statistics of imports of susceptible animals and their products for the past two years, specifying country or zone of origin, species and volume.

6.2.1. Provide a map with the number and location of ports, airports and land crossings. Is the official service responsible for import controls part of the official services, or is it an independent body? If it is an independent body, describe its management structure, staffing levels and resources, and its accountability to the central veterinary services. Describe the communication systems between the central authorities and the border inspection posts, and between border inspection posts.

6.2.2. Provide a description on the methods used for the safe disposal of waste from international traffic, who is responsible and provide a summary, for the past two years, of the quantity disposed of.

6.2.3. Describe the regulations, procedures, type and frequency of checks at the point of entry into the country and/or their final destination, concerning the import and follow up of the following:

a) animals,
b) genetic material (semen and embryos),
c) animal products,
d) veterinary medicinal products (i.e. biologics).

6.2.4. Describe the action available under legislation, and actually taken, when an illegal import is detected. Provide information on detected illegal imports.
7. Control measures and contingency planning

7.1. Give details of any written guidelines, including contingency plans, available to the official services for dealing with suspected or confirmed outbreaks of FMD.

7.2. Is quarantine imposed on premises with suspicious cases, pending final diagnosis? What other procedures are followed regarding suspicious cases?

7.3. In the event of an FMD outbreak:

7.3.1. indicate the sampling and testing procedures used to identify and confirm presence of the causative agent;

7.3.2. describe the actions taken to control the disease situation in and around any holdings found to be infected with FMD;

7.3.3. indicate the control and/or eradication procedures (e.g. vaccination, stamping out, partial slaughter/vaccination etc) that would be taken. Include details on antigen and vaccine banks;

7.3.4. describe the procedures used to confirm that an outbreak has been successfully controlled/eradicated, including any restrictions on restocking;

7.3.5. give details of any compensation payments made available to farmers etc when animals are slaughtered for disease control/eradication purposes and their prescribed timetable.

8. Compliance with the Terrestrial Code

8.1 In addition to the documentary evidence that the provisions of Article 8.5.2. are properly implemented and supervised, the Delegate of the country must submit a declaration indicating:

8.1.1. there has been no outbreak of FMD during the past 12 months;

8.1.2. no evidence of FMDV infection has been found during the past 12 months;

8.1.3. no vaccination against FMD has been carried out during the past 12 months,

8.2. and should confirm that since the cessation of vaccination no animals vaccinated against FMD have been imported.

9. Recovery of status

Countries applying for recovery of status should comply with the provisions of Article 8.5.8. of the Terrestrial Code and provide detailed information as specified in sections 3.1., 3.2., 3.3. and 5.2. of this report. Information in relation to other sections need only be supplied if relevant.
FMD FREE COUNTRY WHERE VACCINATION IS PRACTISED

Report of Country which applies for recognition of status, under Chapter 8.5.
of the Terrestrial Animal Health Code 2008, as a FMD free country practising vaccination

Please address concisely the following topics. National regulations laws and Veterinary Administration directives may be referred to and annexed as appropriate in one of the OIE official languages.

1. Introduction

1.1. Geographical factors. Provide a general description of the country including physical, geographical and other factors that are relevant to FMD dissemination, countries sharing common borders and other countries that although may not be adjacent share a link for the potential introduction of disease. Provide a map identifying the factors above.

1.2. Livestock industry. Provide a general description of the livestock industry in the country.

2. Veterinary system

2.1. Legislation. Provide a list and summary of all relevant veterinary legislation in relation to FMD.

2.2. Veterinary Services. Provide documentation on the compliance of the Veterinary Service of the country with the provisions of Chapters 3.1. and 3.2. of the Terrestrial Code and 1.1.3. of the Terrestrial Manual and describe how the veterinary services supervise and control all FMD related activities. Provide maps and tables wherever possible.

2.3. Role of farmers, industry and other relevant groups in FMD surveillance and control (include a description of training and awareness programs on FMD).

2.4. Role of private veterinary profession in FMD surveillance and control.

3. FMD eradication

3.1. History. Provide a description of the FMD history in the country, date of first detection, origin of infection, date of eradication (date of last case), types and subtypes present.

3.2. Strategy. Describe how FMD was controlled and eradicated (e.g. stamping-out, modified stamping-out, zoning), provide timeframe for eradication.

3.3. Vaccines and vaccination. What type of vaccine is used? What species are vaccinated? Provide evidence that the vaccine used complies with Chapter 2.1.5. of the OIE Terrestrial Manual. Describe the vaccination program, including records kept, and provide evidence to show its effectiveness (e.g. vaccination coverage, serosurveillance, etc.).

3.4. Legislation, organisation and implementation of the FMD eradication campaign. Provide a description of the organizational structure at the different levels. Indicate if detailed operational guidelines exist and give a brief summary.
3.5. Animal identification and movement control. Are susceptible animals identified (individually or at a group level)? Provide a description of the methods of animal identification, herd registration and traceability, including vaccination data. How are animal movements controlled in the country? Provide evidence on the effectiveness of animal identification and movement controls. Please provide information on pastoralism, transhumance and related paths of movement.

4. FMD diagnosis

Provide documentary evidence that the provisions in Chapters 1.1.2., 1.1.3. and 2.1.5. of the Terrestrial Manual are applied. In particular, the following points should be addressed:

4.1. Is FMD laboratory diagnosis carried out in the country? If so, provide a list of approved laboratories. If not, provide the name(s) of and the arrangements with the laboratory (ies) samples are sent to, the follow-up procedures and the time frame for obtaining results.

4.2. Provide an overview of the FMD approved laboratories, in particular to address the following points:

   4.2.1. Procedures for the official accreditation of laboratories. Give details of internal quality management systems, e.g. Good Laboratory Practice, ISO etc. that exist in, or planned for, the laboratory system.

   4.2.2. Give details of participation in inter-laboratory validation tests (ring tests).

   4.2.3. Is live virus handled?

   4.2.4. Biosecurity measures applied.

   4.2.5. Details of the type of tests undertaken.

5. FMD surveillance

Provide documentary evidence that surveillance for FMD in the country complies with the provisions of Articles 8.5.40. to 8.5.46. of the Terrestrial Code and Chapter 2.1.5. of the Terrestrial Manual. In particular, the following points should be addressed:

5.1. Clinical suspicion. What are the criteria for raising a suspicion of FMD? What is the procedure to notify (by whom and to whom) and what penalties are involved for failure to report? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for FMDV, species, type of sample, testing method(s) and results (including differential diagnosis).

5.2. Surveillance. Are serological and virological surveys conducted, in particular applying the provisions of Article 8.5.44? If so, provide detailed information on the survey design (confidence level, sample size, stratification). How frequently are they conducted? Are wildlife susceptible species included in serological surveys? Provide a summary table indicating, for the past two years, the number of samples tested for FMD and FMDV, species, type of sample, testing method(s) and results (including differential diagnosis). Provide details on follow up actions taken on all suspicious and positive results. Provide criteria for selection of populations for targeted surveillance and numbers of animals examined and samples tested. Provide details on the methods applied for monitoring the performance of the surveillance system including indicators.
5.3. Livestock demographics and economics. What is the susceptible animal population by species and production systems? How many herds, flocks, etc., of each susceptible species are in the country? How are they distributed (e.g., herd density, etc.)? Provide tables and maps as appropriate.

5.4. Wildlife demographics. What susceptible species are present in the country? Provide estimates of population sizes and geographic distribution. What are the measures in place to prevent contact between domestic and wildlife susceptible species?

5.5. Slaughterhouses and markets. Where are the major livestock marketing or collection centers? What are the patterns of livestock movement within the country? How are the animals transported and handled during these transactions.

6. FMD prevention

6.1. Coordination with neighbouring countries. Are there any relevant factors about the adjacent countries or zones that should be taken into account (e.g., size, distance from adjacent border to affected herds or animals)? Describe coordination, collaboration and information sharing activities with neighbouring countries.

6.2. Import control procedures

From what countries or zones does the country authorize the import of susceptible animals or their products? What criteria are applied to approve such countries or zones? What controls are applied on entry of such animals and products, and subsequent internal movement? What import conditions and test procedures are required? Are imported animals of susceptible species required to undergo a quarantine or isolation period? If so, for how long and where? Are import permits and health certificates required? What other procedures are used? Provide summary statistics of imports of susceptible animals and their products for the past two years, specifying country or zone of origin, species and volume.

6.2.1. Provide a map with the number and location of ports, airports and land crossings. Is the official service responsible for import controls part of the official services, or is it an independent body? If it is an independent body, describe its management structure, staffing levels and resources, and its accountability to the central veterinary services. Describe the communication systems between the central authorities and the border inspection posts, and between border inspection posts.

6.2.2. Provide a description on the methods used for the safe disposal of waste from international traffic, who is responsible and provide a summary, for the past two years, of the quantity disposed of.

6.2.3. Describe the regulations, procedures, type and frequency of checks at the point of entry into the country and/or their final destination, concerning the import and follow-up of the following:

a) animals,

b) genetic material (semen and embryos),

c) animal products,

d) veterinary medicinal products (i.e. biologics).
6.2.4. Describe the action available under legislation, and actually taken, when an illegal import is detected. Provide information on detected illegal imports.

7. Control measures and contingency planning

7.1. Give details of any written guidelines, including contingency plans, available to the official services for dealing with suspected or confirmed outbreaks of FMD.

7.2. Is quarantine imposed on premises with suspicious cases, pending final diagnosis? What other procedures are followed regarding suspicious cases?

7.3. In the event of an FMD outbreak:

7.3.1. indicate the sampling and testing procedures used to identify and confirm presence of the causative agent;

7.3.2. describe the actions taken to control the disease situation in and around any holdings found to be infected with FMD;

7.3.3. indicate the control and/or eradication procedures (e.g. vaccination, stamping out, partial slaughter/vaccination etc) that would be taken. Include details on antigen and vaccine banks;

7.3.4. describe the procedures used to confirm that an outbreak has been successfully controlled/eradicated, including any restrictions on restocking;

7.3.5. give details of any compensation payments made available to farmers etc when animals are slaughtered for disease control/eradication purposes and their prescribed timetable.

8. Compliance with the Terrestrial Code

8.1. In addition to the documentary evidence that the provisions of Article 8.5.3. are properly implemented and supervised, the Delegate of the country must submit a declaration indicating.

8.2. That there has been no outbreak of FMD for the past 2 years and no evidence of FMDV circulation for the past 12 months, with documented evidence that:

8.2.1. surveillance for FMD and FMDV circulation in accordance with Articles 8.5.40. to 8.5.46. is in operation, and that regulatory measures for the prevention and control of FMD have been implemented;

8.2.2. routine vaccination is carried out for the purpose of the prevention of FMD;

8.2.3. the vaccine used complies with the standards described in the Terrestrial Manual.

9. Recovery of status

Countries applying for recovery of status should comply with the provisions of Article 8.5.8. of the Terrestrial Code and provide detailed information as specified in sections 3.1., 3.2., 3.3. and 5.2. of this report. Information in relation to other sections need only be supplied if relevant.
FMD FREE ZONE WHERE VACCINATION IS NOT PRACTISED

Report of Country which applies for recognition of status, under Chapter 8.5. of the Terrestrial Animal Health Code 2008, for a FMD free zone not practising vaccination

Please address concisely the following topics. National regulations laws and Veterinary Administration directives may be referred to and annexed as appropriate in one of the OIE official languages.

1. Introduction

1.1. Geographical factors. Provide a general description of the country and the zone including physical, geographical and other factors that are relevant to FMD dissemination, countries or zones sharing common borders and other countries or zones that although may not be adjacent share a link for the potential introduction of disease. The boundaries of the zone must be clearly defined, including a buffer zone if applied. Provide a digitalised, geo-referenced map with a precise text description of the geographical boundaries of the zone.

1.2. Livestock industry. Provide a general description of the livestock industry in the country and the zone.

2. Veterinary system

2.1. Legislation. Provide a list and summary of all relevant veterinary legislation in relation to FMD.

2.2. Veterinary Services. Provide documentation on the compliance of the Veterinary Service of the country with the provisions of Chapters 3.1. and 3.2. of the Terrestrial Code and 1.1.3. of the Terrestrial Manual and describe how the veterinary services supervise and control all FMD related activities in the country and in the zone. Provide maps and tables wherever possible.

2.3. Role of farmers, industry and other relevant groups in FMD surveillance and control (include a description of training and awareness programs on FMD).

2.4. Role of private veterinary profession in FMD surveillance and control.

3. FMD eradication

3.1. History. Provide a description of the FMD history in the country and zone, provide date of first detection, origin of infection, date of eradication in the zone (date of last case), types and subtypes present.

3.2. Strategy. Describe how FMD was controlled and eradicated in the zone (e.g. stamping-out, modified stamping-out), provide timeframe for eradication.

3.3. Vaccines and vaccination. If vaccination is used in the rest of the country, what type of vaccine is used? What species are vaccinated? Provide evidence that the vaccine used complies with Chapter 2.1.5. of the Terrestrial Manual. Describe the vaccination program, including records kept, and provide evidence to show its effectiveness (e.g. vaccination coverage, serosurveillance, etc.).
3.4. Legislation, organisation and implementation of the FMD eradication campaign. Provide a description of the organizational structure at the different levels. Indicate if detailed operational guidelines exist and give a brief summary.

3.5. Animal identification and movement control. Are susceptible animals identified (individually or at a group level)? Provide a description of the methods of animal identification, herd registration and traceability. How are animal movements controlled in and between zones of the same or different status, in particular if the provisions of the Terrestrial Code in 8.5.9. are applied? Provide evidence on the effectiveness of animal identification and movement controls. Please provide information on pastoralism, transhumance and the related paths of movement.

4. FMD diagnosis

Provide documentary evidence that the provisions in Chapters 1.1.2., 1.1.3. and 2.1.5. of the Terrestrial Manual are applied. In particular, the following points should be addressed:

4.1. Is FMD laboratory diagnosis carried out in the country? If so, provide a list of approved laboratories. If not, provide the name(s) of and the arrangements with the laboratory (ies) samples are sent to. Indicate the laboratory (ies) where samples originating from the zone are diagnosed, the follow-up procedures and the time frame for obtaining results.

4.2. Provide an overview of the FMD approved laboratories, in particular to address the following points:

4.2.1. Procedures for the official accreditation of laboratories. Give details of internal quality management systems, e.g. Good Laboratory Practice, ISO etc. that exist in, or planned for, the laboratory system.

4.2.2. Give details of participation in inter-laboratory validation tests (ring tests).

4.2.3. Is live virus handled?

4.2.4. Biosecurity measures applied.

4.2.5. Details of the type of tests undertaken.

5. FMD surveillance

Provide documentary evidence that surveillance for FMD in the zone complies with the provisions of Articles 8.5.40. to 8.5.46. of the Terrestrial Code and Chapter 2.1.5. of the Terrestrial Manual. In particular, the following points should be addressed:

5.1. Clinical suspicion. What are the criteria for raising a suspicion of FMD? What is the procedure to notify (by whom and to whom) and what penalties are involved for failure to report? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for FMDV, species, type of sample, testing method(s) and results (including differential diagnosis).
5.2. Serological surveillance. Are serological surveys conducted? If so, provide detailed information on the survey design (confidence level, sample size, stratification). How frequently are they conducted? Are wildlife susceptible species included in serological surveys? Provide a summary table indicating, for the past two years, the number of samples tested for FMDV, species, type of sample, testing method(s) and results (including differential diagnosis). Provide details on follow up actions taken on all suspicious and positive results. Provide criteria for selection of populations for targeted surveillance and numbers of animals examined and samples tested. Provide details on the methods applied for monitoring the performance of the surveillance system including indicators.

5.3. Livestock demographics and economics. What is the susceptible animal population by species and production systems in the country and the zone? How many herds, flocks, etc., of each susceptible species are in the country? How are they distributed (e.g., herd density, etc.)? Provide tables and maps as appropriate.

5.4. Wildlife demographics. What susceptible species are present in the country and the zone? Provide estimates of population sizes and geographic distribution. What are the measures in place to prevent contact between domestic and wildlife susceptible species?

5.5. Slaughterhouses and markets. Where are the major livestock marketing or collection centers? What are the patterns of livestock movement within the country? How are the animals transported and handled during these transactions.

6. FMD prevention

6.1. Coordination with neighbouring countries. Are there any relevant factors about the adjacent countries and zones that should be taken into account (e.g., size, distance from adjacent border to affected herds or animals)? Describe coordination, collaboration and information sharing activities with neighbouring countries and zones.

If the FMD free zone without vaccination is situated in an FMD infected country or borders and infected country or zone, describe the animal health measures implemented to effectively prevent the introduction of the agent, taking into consideration physical or geographical barriers.

6.2. Import control procedures

From what countries or zones does the country authorize the import of susceptible animals or their products into free zone? What criteria are applied to approve such countries or zones? What controls are applied on entry of such animals and products, and subsequent internal movement? What import conditions and test procedures are required? Are imported animals of susceptible species required to undergo a quarantine or isolation period? If so, for how long and where? Are import permits and health certificates required? What other procedures are used? Provide summary statistics of imports of susceptible animals and their products for the past two years, specifying country or zone of origin, species and volume.

6.2.1. Provide a map with the number and location of ports, airports and land crossings. Is the official service responsible for import controls part of the official services, or is it an independent body? If it is an independent body, describe its management structure, staffing levels and resources, and its accountability to the central veterinary services. Describe the communication systems between the central authorities and the border inspection posts, and between border inspection posts.
6.2.2. Provide a description on the methods used for the safe disposal of waste from international traffic, who is responsible and provide a summary, for the past two years, of the quantity disposed of.

6.2.3. Describe the regulations, procedures, type and frequency of checks at the point of entry into the country and/or their final destination, concerning the import and follow up of the following:

a) animals,
b) genetic material (semen and embryos),
c) animal products,
d) veterinary medicinal products (i.e. biologics).

6.2.4. Describe the action available under legislation, and actually taken, when an illegal import is detected. Provide information on detected illegal imports.

7. Control measures and contingency planning

7.1. Give details of any written guidelines, including contingency plans, available to the official services for dealing with suspected or confirmed outbreaks of FMD.

7.2. Is quarantine imposed on premises with suspicious cases, pending final diagnosis? What other procedures are followed regarding suspicious cases?

7.3. In the event of an FMD outbreak:

7.3.1. indicate the sampling and testing procedures used to identify and confirm presence of the causative agent;

7.3.2. describe the actions taken to control the disease situation in and around any holdings found to be infected with FMD;

7.3.3. indicate the control and/or eradication procedures (eg. vaccination, stamping out, partial slaughter/vaccination etc) that would be taken. Include details on antigen and vaccine banks;

7.3.4. describe the procedures used to confirm that an outbreak has been successfully controlled/eradicated, including any restrictions on restocking;

7.3.5. give details of any compensation payments made available to farmers etc when animals are slaughtered for disease control/eradication purposes and their prescribed timetable.

8. Compliance with the Terrestrial Code

8.1. In addition to the documentary evidence that the provisions of Article 8.5.4. are properly implemented and supervised, the Delegate of the country must submit a declaration indicating:

8.1.1. there has been no outbreak of FMD during the past 12 months;

8.1.2. no evidence of FMDV infection has been found during the past 12 months;
8.1.3. no vaccination against FMD has been carried out during the past 12 months;

8.1.4. no vaccinated animal has been introduced into the zone since the cessation of vaccination, except in accordance with Article 8.5.9.

9. Recovery of status

Countries applying for recovery of status should comply with the provisions of Article 8.5.8. of the Terrestrial Code and provide detailed information as specified in sections 3.1., 3.2., 3.3. and 5.2. of this report. Information in relation to other sections need only be supplied if relevant.

FMD FREE ZONE WHERE VACCINATION IS PRACTISED

Report of Country which applies for recognition of status, under Chapter 8.5. of the Terrestrial Animal Health Code 2008, for a FMD free zone practising vaccination

Please address concisely the following topics. National regulations laws and Veterinary Administration directives may be referred to and annexed as appropriate in one of the OIE official languages.

1. Introduction

1.1. Geographical factors. Provide a general description of the country and the zone including physical, geographical and other factors that are relevant to FMD dissemination, countries or zones sharing common borders and other countries or zones that although may not be adjacent share a link for the potential introduction of disease. The boundaries of the zone must be clearly defined, including a buffer zone if applied. Provide a digitalised, geo-referenced map with a precise text description of the geographical boundaries of the zone.

1.2. Livestock industry. Provide a general description of the livestock industry in the country and the zone.

2. Veterinary system

2.1. Legislation. Provide a list and summary of all relevant veterinary legislation in relation to FMD.

2.2. Veterinary Services. Provide documentation on the compliance of the Veterinary Service of the country with the provisions of Chapters 3.1. and 3.2. of the Terrestrial Code and 1.1.3. of the Terrestrial Manual and describe how the veterinary services supervise and control all FMD related activities in the country and in the zone. Provide maps and tables wherever possible.

2.3. Role of farmers, industry and other relevant groups in FMD surveillance and control (include a description of training and awareness programs on FMD).

2.4. Role of private veterinary profession in FMD surveillance and control.
3. FMD eradication

3.1. History. Provide a description of the FMD history in the country and zone, provide date of first detection, origin of infection, date of eradication in the zone (date of last case), types and subtypes present.

3.2. Strategy. Describe how FMD was controlled and eradicated in the zone (e.g. stamping-out, modified stamping-out), provide timeframe for eradication.

3.3. Vaccines and vaccination. What type of vaccine is used? What species are vaccinated? Provide evidence that the vaccine used complies with Chapter 2.1.5. of the Terrestrial Manual. Describe the vaccination program in the country and in the zone, including records kept, and provide evidence to show its effectiveness (e.g. vaccination coverage, serosurveillance, etc.).

3.4. Legislation, organisation and implementation of the FMD eradication campaign. Provide a description of the organizational structure at the different levels. Indicate if detailed operational guidelines exist and give a brief summary.

3.5. Animal identification and movement control. Are susceptible animals identified (individually or at a group level)? Provide a description of the methods of animal identification, herd registration and traceability, including vaccination data. How are animal movements controlled in and between zones of the same or different status, in particular if the provisions of the Terrestrial Code in 8.5.9. are applied? Provide evidence on the effectiveness of animal identification and movement controls. Please provide information on pastoralism, transhumance and the related paths of movement.

4. FMD diagnosis

Provide documentary evidence that the provisions in Chapters 1.1.2., 1.1.3. and 2.1.5. of the Terrestrial Manual are applied. In particular, the following points should be addressed:

4.1. Is FMD laboratory diagnosis carried out in the country? If so, provide a list of approved laboratories. If not, provide the name(s) of and the arrangements with the laboratory (ies) samples are sent to, the follow up procedures and the time frame for obtaining results. Indicate the laboratory (ies) where samples originating from the zone are diagnosed.

4.2. Provide an overview of the FMD approved laboratories, in particular to address the following points:

4.2.1. Procedures for the official accreditation of laboratories. Give details of internal quality management systems, e.g. Good Laboratory Practice, ISO etc. that exist in, or planned for, the laboratory system.

4.2.2. Give details of participation in inter-laboratory validation tests (ring tests).

4.2.3. Is live virus handled?

4.2.4. Biosecurity measures applied.

4.2.5. Details of the type of tests undertaken.
5. FMD surveillance

Provide documentary evidence that surveillance for FMD in the zone complies with the provisions of Articles 8.5.40. to 8.5.46. of the Terrestrial Code and Chapter 2.1.5. of the Terrestrial Manual. In particular, the following points should be addressed:

5.1 Clinical suspicion. What are the criteria for raising a suspicion of FMD? What is the procedure to notify (by whom and to whom) and what penalties are involved for failure to report? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for FMDV, species, type of sample, testing method(s) and results (including differential diagnosis).

5.2 Surveillance. Are serological and virological surveys conducted, in particular applying the provisions of Article 8.5.44.? If so, provide detailed information on the survey design (confidence level, sample size, stratification). How frequently are they conducted? Are wildlife susceptible species included in serological surveys? Provide a summary table indicating, for the past two years, the number of samples tested for FMD and FMDV, species, type of sample, testing method(s) and results (including differential diagnosis). Provide details on follow up actions taken on all suspicious and positive results. Provide criteria for selection of populations for targeted surveillance and numbers of animals examined and samples tested. Provide details on the methods applied for monitoring the performance of the surveillance system including indicators.

5.3 Livestock demographics and economics. What is the susceptible animal population by species and production systems in the country and the zone? How many herds, flocks, etc., of each susceptible species are in the country? How are they distributed (e.g., herd density, etc.)? Provide tables and maps as appropriate.

5.4 Wildlife demographics. What susceptible species are present in the country and in the zone? Provide estimates of population sizes and geographic distribution. What are the measures in place to prevent contact between domestic and wildlife susceptible species?

5.5 Slaughterhouses and markets. Where are the major livestock marketing or collection centers? What are the patterns of livestock movement within the country? How are the animals transported and handled during these transactions.

6. FMD prevention

6.1 Coordination with neighbouring countries. Are there any relevant factors about the adjacent countries and zones that should be taken into account (e.g. size, distance from adjacent border to affected herds or animals)? Describe coordination, collaboration and information sharing activities with neighbouring countries and zones.

If the FMD free zone with vaccination is situated in an FMD infected country or borders and infected country or zone, describe the animal health measures implemented to effectively prevent the introduction of the agent, taking into consideration physical or geographical barriers.
6.2. Import control procedures

From what countries or zones does the country authorize the import of susceptible animals or their products into free zone? What criteria are applied to approve such countries or zones? What controls are applied on entry of such animals and products, and subsequent internal movement? What import conditions and test procedures are required? Are imported animals of susceptible species required to undergo a quarantine or isolation period? If so, for how long and where? Are import permits and health certificates required? What other procedures are used? Provide summary statistics of imports of susceptible animals and their products for the past two years, specifying country or zone of origin, species and volume.

6.2.1. Provide a map with the number and location of ports, airports and land crossings. Is the official service responsible for import controls part of the official services, or is it an independent body? If it is an independent body, describe its management structure, staffing levels and resources, and its accountability to the central veterinary services. Describe the communication systems between the central authorities and the border inspection posts, and between border inspection posts.

6.2.2. Provide a description on the methods used for the safe disposal of waste from international traffic, who is responsible and provide a summary, for the past two years, of the quantity disposed of.

6.2.3. Describe the regulations, procedures, type and frequency of checks at the point of entry into the country and/or their final destination, concerning the import and follow up of the following:

a) animals,
b) genetic material (semen and embryos),
c) animal products,
d) veterinary medicinal products (i.e. biologics).

6.2.4. Describe the action available under legislation, and actually taken, when an illegal import is detected. Provide information on detected illegal imports.

7. Control measures and contingency planning

7.1. Give details of any written guidelines, including contingency plans, available to the official services for dealing with suspected or confirmed outbreaks of FMD.

7.2. Is quarantine imposed on premises with suspicious cases, pending final diagnosis? What other procedures are followed regarding suspicious cases?

7.3. In the event of an FMD outbreak:

7.3.1. indicate the sampling and testing procedures used to identify and confirm presence of the causative agent;

7.3.2. describe the actions taken to control the disease situation in and around any holdings found to be infected with FMD;
Annex XXXII (contd)

7.3.3. indicate the control and/or eradication procedures (e.g. vaccination, stamping-out, partial slaughter/vaccination etc) that would be taken. Include details on antigen and vaccine banks;

7.3.4. describe the procedures used to confirm that an outbreak has been successfully controlled/eradicated, including any restrictions on restocking.

7.3.5. Give details of any compensation payments made available to farmers etc when animals are slaughtered for disease control/eradication purposes and their prescribed timetable.

8. **Compliance with the Terrestrial Code**

8.1. In addition to the documentary evidence that the provisions of Article 8.5.5. are properly implemented and supervised, the Delegate of the country must submit a declaration indicating:

8.1.1. that there has been no outbreak of FMD for the past 2 years,

8.1.2. no evidence of FMDV circulation for the past 12 months,

8.1.3. surveillance for FMD and FMDV circulation in accordance with Articles 8.5.40. to 8.5.46. is in operation.

9. **Recovery of status**

Countries applying for recovery of status should comply with the provisions of Article 8.5.8. of the Terrestrial Code and provide detailed information as specified in sections 3.1., 3.2., 3.3. and 5.2. of this report. Information in relation to other sections need only be supplied if relevant.

**Article 4.**

**RINDERPEST QUESTIONNAIRE**

**RINDERPEST INFECTION FREE COUNTRY**


Please address concisely the following topics. National regulations laws and Veterinary Administration directives may be referred to and annexed as appropriate in one of the OIE official languages.

1. **Introduction**

1.1. Geographical factors. Provide a general description of the country including physical, geographical and other factors that are relevant to rinderpest dissemination, countries sharing common borders and other countries that although may not be adjacent share a link for the potential introduction of disease. Provide a map identifying the factors above.

1.2. Livestock industry. Provide a general description of the livestock industry in the country.
2. Veterinary system

2.1. Legislation. Provide a list and summary of all relevant veterinary legislation in relation to rinderpest.

2.2. Veterinary Services. Provide documentation on the compliance of the Veterinary Service of the country with the provisions of chapters 3.1. and 3.2. of the Terrestrial Code and 1.1.3. of the Terrestrial Manual and describe how the veterinary services supervise and control all rinderpest related activities. Provide maps and tables wherever possible.

2.3. Role of farmers, industry and other relevant groups in rinderpest surveillance and control (include a description of training and awareness programs on rinderpest).

2.4. Role of private veterinary profession in rinderpest surveillance and control.

3. Rinderpest eradication

3.1. History. Provide a description of the rinderpest history in the country, date of first detection, origin of infection, date of eradication (date of last case), lineage(s) present.

3.2. Strategy. Describe how rinderpest was controlled and eradicated (e.g. stamping-out, modified stamping-out, zoning), provide timeframe for eradication.

3.3. Vaccines and vaccination. Was rinderpest vaccine ever used? If so, when was the last vaccination carried out? What species were vaccinated? Has heterologous vaccine been used in cattle, buffalo or yak?

3.4. Legislation, organisation and implementation of the rinderpest eradication campaign. Provide a description of the organizational structure at the different levels. Indicate if detailed operational guidelines exist and give a brief summary.

3.5. Animal identification and movement control. Are susceptible animals identified (individually or at a group level)? Provide a description of the methods of animal identification, herd registration and traceability. How are animal movements controlled in the country? Provide evidence on the effectiveness of animal identification and movement controls. Please provide information on pastoralism, transhumance and related paths of movement.

4. Rinderpest diagnosis

Provide evidence that a system is in place for the rapid confirmation of a suspected outbreak i.e. that the provisions in Chapters 1.1.3. and 2.1.15. of the Terrestrial Manual are applied. In particular, the following points should be addressed:

4.1. Is rinderpest laboratory diagnosis carried out in the country? If so, provide a list of approved laboratories. If not, provide the name(s) of and the arrangements with the laboratory(ies) samples are sent to, the follow up procedures and the timeframe for obtaining results.

4.2. Provide an overview of the rinderpest approved laboratories, in particular to address the following points:
4.2.1. Procedures for the official accreditation of laboratories. Give details of internal quality management systems, e.g. Good Laboratory Practice, ISO etc. that exist in, or planned for, the laboratory system.

4.2.2. Give details of participation in inter-laboratory validation tests (ring tests).

4.2.3. Is live virus handled?

4.2.4. Biosecurity measures applied.

4.2.5. Details of the type of tests undertaken.

5. Rinderpest surveillance

Provide documentary evidence that surveillance for rinderpest in the country complies with the provisions of Articles 8.13.20. to 8.13.27. of the Terrestrial Code and Chapter 2.1.15. of the Terrestrial Manual. In particular, the following points should be addressed:

5.1. Clinical suspicion. What are the criteria for raising a suspicion of rinderpest? What is the procedure to notify (by whom and to whom) and what penalties are involved for failure to report? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for rinderpest virus, species, type of sample, testing method(s) and results (including differential diagnosis). In particular, provide evidence of compliance with the provisions of Articles 8.13.20. to 8.13.27. of the Terrestrial Code.

5.2. Serological surveillance. Are serological surveys conducted? If so, provide detailed information on the survey design in accordance with Articles 8.13.20. to 8.13.27. of the Terrestrial Code. Are wildlife susceptible species included in serological surveys? If not, explain the rationale. Provide a summary table indicating, for the past two years, the number of samples tested for rinderpest virus, species, type of sample, testing method(s) and results (including differential diagnosis). Provide details on follow up actions taken on all suspicious and positive results. Provide criteria for selection of populations for targeted surveillance and numbers of animals examined and samples tested. Provide details on the methods applied for monitoring the performance of the surveillance system including indicators.

5.3. Livestock demographics and economics. What is the susceptible animal population by species and production systems? How many herds, flocks, etc., of each susceptible species are in the country? How are they distributed (e.g. herd density, etc.)? Provide tables and maps as appropriate.

5.4. Wildlife demographics. What susceptible species are present in the country? Provide estimates of population sizes and geographic distribution. What are the measures in place to prevent contact between domestic and wildlife susceptible species?

5.5. Slaughterhouses and markets. Where are the major livestock marketing or collection centers? What are the patterns of livestock movement within the country? How are the animals transported and handled during these transactions.

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Accounts of the ages for eruption of the incisor teeth vary markedly and are clearly dependent on species, breed, nutritional status and nature of the feed. Therefore, for the purposes of serosurveillance, it should be noted that:

a) cattle having only one pair of erupted permanent central incisor teeth are aged between 21 and 36 months (Asian buffalos 24-48 months);

b) cattle having only two pairs of erupted permanent central incisor teeth are aged between 30 and 48 months (Asian buffalos 48-60 months).
6. **Rinderpest prevention**

6.1. Coordination with neighbouring countries. Are there any relevant factors about the adjacent countries that should be taken into account (e.g., size, distance from adjacent border to affected herds or animals)? Describe coordination, collaboration and information sharing activities with neighbouring countries.

6.2. Import control procedures

From what countries or zones does the country authorize the import of susceptible animals or their products? What criteria are applied to approve such countries or zones? What controls are applied on entry of such animals and products, and subsequent internal movement? What import conditions and test procedures are required? Are imported animals of susceptible species required to undergo a quarantine or isolation period? If so, for how long and where? Are import permits and health certificates required? What other procedures are used? Provide summary statistics of imports of susceptible animals and their products for the past two years, specifying country or zone of origin, species and volume.

6.2.1. Provide a map with the number and location of ports, airports and land crossings. Is the official service responsible for import controls part of the official services, or is it an independent body? If it is an independent body, describe its management structure, staffing levels and resources, and its accountability to the central veterinary services. Describe the communication systems between the central authorities and the border inspection posts, and between border inspection posts.

6.2.2. Describe the regulations, procedures, type and frequency of checks at the point of entry into the country and/or their final destination, concerning the import and follow up of the following:

   a) animals,

   b) genetic material (semen and embryos),

   c) animal products,

   d) veterinary medicinal products (i.e. biologics).

6.2.3. Describe the action available under legislation, and actually taken, when an illegal import is detected. Provide information on detected illegal imports.

7. **Control measures and contingency planning**

7.1. Give details of any written guidelines, including contingency plans, available to the official services for dealing with suspected or confirmed outbreaks of rinderpest.

7.2. Is quarantine imposed on premises with suspicious cases, pending final diagnosis? What other procedures are followed regarding suspicious cases?
7.3. In the event of a rinderpest outbreak:

7.3.1 indicate the sampling and testing procedures used to identify and confirm presence of the causative agent;

7.3.2 describe the actions taken to control the disease situation in and around any holdings found to be infected with rinderpest;

7.3.3 indicate the control and/or eradication procedures (e.g. vaccination, stamping out, partial slaughter/vaccination etc) that would be taken;

7.3.4 describe the procedures used to confirm that an outbreak has been successfully controlled/eradicated, including any restrictions on restocking;

7.3.5 give details of any compensation payments made available to farmers etc when animals are slaughtered for disease control/eradication purposes and their prescribed timetable.

8. Compliance with the Terrestrial Code

8.1. The Delegate of the country must submit documentary evidence that the provisions of Article 8.13.2. or 1.4.6.1. (historical freedom) of the Terrestrial Code have been properly implemented and supervised.

9. Recovery of status

Countries applying for recovery of status should comply with the provisions of Article 8.13.3. of the Terrestrial Code and provide detailed information as specified in sections 3.1., 3.2., 3.3. and 5.2. of this questionnaire. Information in relation to other sections need only be supplied if relevant.

Article 5.

CBPP QUESTIONNAIRE

CBPP FREE COUNTRY

Report of Country which applies for recognition of status, under Chapter 2.3.15 and Appendix 3.8.3 of the Terrestrial Animal Health Code, as a CBPP free country

Please address concisely the following topics. National regulations laws and Veterinary Administration directives may be referred to and annexed as appropriate in one of the OIE official languages.

1. Introduction

1.1. Geographical factors. Provide a general description of the country including physical, geographical and other factors that are relevant to CBPP dissemination, countries sharing common borders and other countries that although may not be adjacent share a link for the potential introduction of disease. Provide a map identifying the factors above.

1.2. Livestock industry. Provide a general description of the livestock industry in the country.
2. Veterinary system

2.1. Legislation. Provide a list and summary of all relevant veterinary legislation in relation to CBPP.

2.2. Veterinary Services. Provide documentation on the compliance of the Veterinary Service of the country with the provisions of Chapters 1.3.3. and 1.3.4. of the Terrestrial Code and I.1.2 of the Terrestrial Manual and describe how the veterinary services supervise and control all CBPP related activities. Provide maps and tables wherever possible.

2.3. Role of farmers, industry and other relevant groups in CBPP surveillance and control (include a description of training and awareness programs on CBPP).

2.4. Role of private veterinary profession in CBPP surveillance and control.

3. CBPP eradication

3.1. History. Provide a description of the CBPP history in the country, date of first detection, origin of infection, date of eradication (date of last case).

3.2. Strategy. Describe how CBPP was controlled and eradicated (e.g. stamping-out, modified stamping-out, zoning), provide timeframe for eradication.

3.3. Vaccines and vaccination. Was CBPP vaccine ever used? If so, when was the last vaccination carried out?

3.4. Legislation, organisation and implementation of the CBPP eradication campaign. Provide a description of the organizational structure at the different levels. Indicate if detailed operational guidelines exist and give a brief summary.

3.5. Animal identification and movement control. Are susceptible animals identified (individually or at a group level)? Provide a description of the methods of animal identification, herd registration and traceability. How are animal movements controlled in the country? Provide evidence on the effectiveness of animal identification and movement controls. Please provide information on pastoralism, transhumance and the related paths of movement.

4. CBPP diagnosis

Provide documentary evidence that the provisions in Chapters I.1.2 and 2.1.6. of the Terrestrial Manual are applied. In particular, the following points should be addressed:

4.1. Is CBPP laboratory diagnosis carried out in the country? If so, provide a list of approved laboratories. If not, provide the name(s) of and the arrangements with the laboratory(ies) samples are sent to, the follow up procedures and the time frame for obtaining results.

4.2. Provide an overview of the CBPP approved laboratories, in particular to address the following points:

4.2.1. Procedures for the official accreditation of laboratories. Give details of internal quality management systems, e.g. Good Laboratory Practice, ISO etc. that exist in, or planned for, the laboratory system.
Annex XXXII (contd)

4.2.2. Give details of participation in inter-laboratory validation tests (ring tests).

4.2.3. Biosecurity measures applied.

4.2.4. Details of the type of tests undertaken including procedures to isolate and identify M. mycoides subsp. mycoides SC as opposed to M. mycoides subsp. mycoides LC.

5. CBPP surveillance

Provide documentary evidence that surveillance for CBPP in the country complies with the provisions of Article 11.8.14. of the Terrestrial Code and Chapter 2.1.6 of the Terrestrial Manual. In particular, the following points should be addressed:

5.1. Clinical surveillance. What are the criteria for raising a suspicion of CBPP? What is the procedure to notify (by whom and to whom) and what penalties are involved for failure to report? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for CBPP agent, species, type of sample, testing method(s) and results (including differential diagnosis).

5.2. Slaughterhouses, slaughter slabs, abattoirs. What are the criteria for raising a suspicion of CBPP lesion? What is the procedure to notify (by whom and to whom)? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for CBPP agent, species, type of sample, testing method(s) and results (including differential diagnosis).

5.3. Provide details on training programs for personnel involved in clinical and slaughter facilities surveillance, and the approaches used to increase community involvement in CBPP surveillance programs.

5.4. For countries where a significant proportion of animals are not slaughtered in controlled abattoirs, what are the alternative surveillance measures applied to detect CBPP (e.g. active clinical surveillance programs, laboratory follow up).

5.5. Livestock demographics and economics. What is the susceptible animal population by species and production systems? How many herds of each susceptible species are in the country? How are they distributed (e.g., herd density, etc.)? Provide tables and maps as appropriate.

5.6. Slaughterhouses and markets. Where are the major livestock marketing or collection centers? What are the patterns of livestock movement within the country? How are the animals transported and handled during these transactions.

5.7. Provide a description of the means employed during the two years preceding this application to rule out the presence of any MmmSC strain in the susceptible population. Provide criteria for selection of populations for targeted surveillance and numbers of animals examined and samples tested. Provide details on the methods applied for monitoring the performance of the surveillance system including indicators.
6. **CBPP prevention**

6.1. Coordination with neighbouring countries. Are there any relevant factors about the adjacent countries that should be taken into account (e.g., size, distance from adjacent border to affected herds or animals)? Describe coordination, collaboration and information sharing activities with neighboring countries.

6.2. Import control procedures

From what countries or zones does the country authorize the import of susceptible animals? What criteria are applied to approve such countries or zones? What controls are applied on entry of such animals, and subsequent internal movement? What import conditions and test procedures are required? Are imported animals of susceptible species required to undergo a quarantine or isolation period? If so, for how long and where? Are import permits and health certificates required? What other procedures are used? Provide summary statistics of imports of susceptible animals for the past two years, specifying country or zone of origin, species and volume.

6.2.1. Provide a map with the number and location of ports, airports and land crossings. Is the official service responsible for import controls part of the official services, or is it an independent body? If it is an independent body, describe its management structure, staffing levels and resources, and its accountability to the central veterinary services. Describe the communication systems between the central authorities and the border inspection posts, and between border inspection posts.

6.2.2. Describe the regulations, procedures, type and frequency of checks at the point of entry into the country and/or their final destination, concerning the import and follow up of the following:

- a) animals,
- b) veterinary medicinal products (i.e. biologics).

6.2.3. Describe the action available under legislation, and actually taken, when an illegal import is detected. Provide information on detected illegal imports.

7. **Control measures and contingency planning**

7.1. Give details of any written guidelines, including contingency plans, available to the official services for dealing with suspected or confirmed outbreaks of CBPP.

7.2. Is quarantine imposed on premises with suspicious cases, pending final diagnosis? What other procedures are followed regarding suspicious cases?

7.3. In the event of a CBPP outbreak:

- 7.3.1. indicate the sampling and testing procedures used to identify and confirm presence of the causative agent;
- 7.3.2. describe the actions taken to control the disease situation in and around any holdings found to be infected with CBPP;
Annex XXXII (contd)

7.3.3. indicate the control and/or eradication procedures (e.g. vaccination, stamping out, partial slaughter/vaccination etc.) that would be taken;

7.3.4. describe the procedures used to confirm that an outbreak has been successfully controlled/eradicated, including any restrictions on restocking;

7.3.5. give details of any compensation payments made available to farmers etc when animals are slaughtered for disease control/eradication purposes and their prescribed timetable.

8. Compliance with the Terrestrial Code

In addition to the documentary evidence that the provisions of Article 11.8.14. are properly implemented and supervised, the Delegate of the country must submit a declaration indicating:

8.1. no clinical CBPP has been detected for at least 2 years;

8.2. no CBPP vaccines have been used for at least 2 years in any susceptible species;

8.3. the country operates both clinical surveillance and disease reporting systems for CBPP adequate to detect clinical disease if it were present;

8.4. all clinical and pathological evidence suggestive of CBPP is investigated by field and laboratory methods (including serological assessment) to refute a possible diagnosis of CBPP;

8.5. there are effective measures in force to prevent the re-introduction of the disease.

9. Recovery of status

Countries applying for recovery of status should comply with the provisions of Article 2.3.15.3 of the Terrestrial Code and provide detailed information as specified in sections 3.1, 3.2, 3.3, 5.2, 5.3 and 5.4 of this report. Information in relation to other sections need only be supplied if relevant.

CBPP FREE ZONE


Please address concisely the following topics. National regulations laws and Veterinary Administration directives may be referred to and annexed as appropriate in one of the OIE official languages.

1. Introduction

1.1. Geographical factors. Provide a general description of the country including physical, geographical and other factors that are relevant to CBPP dissemination, countries sharing common borders and other countries that although may not be adjacent share a link for the potential introduction of disease. Provide a map identifying the factors above. The boundaries of the zone must be clearly defined. Provide a digitalised, geo-referenced map with a precise text description of the geographical boundaries of the zone.

1.2. Livestock industry. Provide a general description of the livestock industry in the country.
2. **Veterinary system**

2.1. Legislation. Provide a list and summary of all relevant veterinary legislation in relation to CBPP.

2.2. Veterinary Services. Provide documentation on the compliance of the Veterinary Service of the country with the provisions of Chapters 1.3.3. and 1.3.4. of the Terrestrial Code and I.1.2 of the Terrestrial Manual and describe how the veterinary services supervise and control all CBPP related activities. Provide maps and tables wherever possible.

2.3. Role of farmers, industry and other relevant groups in CBPP surveillance and control (include a description of training and awareness programs on CBPP).

2.4. Role of private veterinary profession in CBPP surveillance and control.

3. **CBPP eradication**

3.1. History. Provide a description of the CBPP history in the country, date of first detection, origin of infection, date of eradication (date of last case).

3.2. Strategy. Describe how CBPP was controlled and eradicated in the zone (e.g. stamping-out, modified stamping-out, zoning), provide timeframe for eradication.

3.3. Vaccines and vaccination. Was CBPP vaccine ever used? In the entire country? If vaccination was used, when was the last vaccination carried out? Where in the country?

3.4. Legislation, organisation and implementation of the CBPP eradication campaign. Provide a description of the organizational structure at the different levels. Indicate if detailed operational guidelines exist and give a brief summary.

3.5. Animal identification and movement control. Are susceptible animals identified (individually or at a group level)? Provide a description of the methods of animal identification, herd registration and traceability. How are animal movements controlled in the zone? Provide evidence on the effectiveness of animal identification and movement controls. Please provide information on pastoralism, transhumance and the related paths of movement.

4. **CBPP diagnosis**

Provide documentary evidence that the provisions in Chapters I.1.2 and 2.1.6. of the Terrestrial Manual are applied. In particular, the following points should be addressed:

4.1. Is CBPP laboratory diagnosis carried out in the country? If so, provide a list of approved laboratories. If not, provide the name(s) of and the arrangements with the laboratory(ies) samples are sent to, the follow up procedures and the time frame for obtaining results.

4.2. Provide an overview of the CBPP approved laboratories, in particular to address the following points:

4.2.1. Procedures for the official accreditation of laboratories. Give details of internal quality management systems, e.g. Good Laboratory Practice, ISO, etc. that exist in, or planned for, the laboratory system.
4.2.2. Give details of participation in inter-laboratory validation tests (ring tests).

4.2.3. Biosecurity measures applied.

4.2.4. Details of the type of tests undertaken including procedures to isolate and identify M. mycoides subsp. mycoides SC as opposed to M. mycoides subsp. mycoides LC.

5. CBPP surveillance

Provide documentary evidence that surveillance for CBPP in the country complies with the provisions of Article 11.8.14. of the Terrestrial Code and Chapter 2.1.6. of the Terrestrial Manual. In particular, the following points should be addressed:

5.1. Clinical surveillance. What are the criteria for raising a suspicion of CBPP? What is the procedure to notify (by whom and to whom) and what penalties are involved for failure to report? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for CBPP agent, species, type of sample, testing method(s) and results (including differential diagnosis).

5.2. Slaughterhouses, slaughter slabs, abattoirs. What are the criteria for raising a suspicion of CBPP lesion? What is the procedure to notify (by whom and to whom)? Provide a summary table indicating, for the past two years, the number of suspect cases, the number of samples tested for CBPP agent, species, type of sample, testing method(s) and results (including differential diagnosis).

5.3. Provide details on training programs for personnel involved in clinical and slaughter facilities surveillance, and the approaches used to increase community involvement in CBPP surveillance programs.

5.4. For countries where a significant proportion of animals in the zone are not slaughtered in controlled abattoirs, what are the alternative surveillance measures applied to detect CBPP (e.g. active clinical surveillance program, laboratory follow up).

5.5. Livestock demographics and economics. What is the susceptible animal population by species and production systems? How many herds of each susceptible species are in the zone? How are they distributed (e.g., herd density, etc.)? Provide tables and maps as appropriate.

5.6. Slaughterhouses and markets. Where are the major livestock marketing or collection centers? What are the patterns of livestock movement within the country and the zone? How are the animals transported and handled during these transactions.

5.7. Provide a description of the means employed during the two years preceding this application to rule out the presence of any MmmSC strain in the susceptible population of the zone. Provide criteria for selection of populations for targeted surveillance and numbers of animals examined and samples tested. Provide details on the methods applied for monitoring the performance of the surveillance system including indicators.
6. CBPP prevention

6.1. Coordination with neighbouring countries and zones. Are there any relevant factors about the adjacent countries and zones that should be taken into account (e.g. size, distance from adjacent border to affected herds or animals)? Describe coordination, collaboration and information sharing activities with neighboring countries and zones. If the CBPP free zone is situated in a CBPP infected country or borders an infected country or zone, describe the animal health measures implemented to effectively prevent the introduction of the agent, taking into consideration physical or geographical barriers.

6.2. Import control procedures

From what countries or zones does the country authorize the import of susceptible animals? What criteria are applied to approve such countries or zones? What controls are applied on entry of such animals, and subsequent internal movement? What import conditions and test procedures are required? Are imported animals of susceptible species required to undergo a quarantine or isolation period? If so, for how long and where? Are import permits and health certificates required? What other procedures are used? Provide summary statistics of imports of susceptible animals for the past two years, specifying country or zone of origin, species and volume.

6.2.1. Provide a map with the number and location of ports, airports and land crossings. Is the official service responsible for import controls part of the official services, or is it an independent body? If it is an independent body, describe its management structure, staffing levels and resources, and its accountability to the central veterinary services. Describe the communication systems between the central authorities and the border inspection posts, and between border inspection posts.

6.2.2. Describe the regulations, procedures, type and frequency of checks at the point of entry into the zone and/or their final destination, concerning the import and follow up of the following:

a) animals,

b) veterinary medicinal products (i.e. biologics).

6.2.3. Describe the action available under legislation, and actually taken, when an illegal import is detected. Provide information on detected illegal imports.

7. Control measures and contingency planning

7.1. Give details of any written guidelines, including contingency plans, available to the official services for dealing with suspected or confirmed outbreaks of CBPP.

7.2. Is quarantine imposed on premises with suspicious cases, pending final diagnosis? What other procedures are followed regarding suspicious cases?

7.3. In the event of a CBPP outbreak:

7.3.1. indicate the sampling and testing procedures used to identify and confirm presence of the causative agent;

7.3.2. describe the actions taken to control the disease situation in and around any holdings found to be infected with CBPP;
7.3.3. indicate the control and/or eradication procedures (e.g. vaccination, stamping out, partial slaughter/vaccination, etc.) that would be taken;

7.3.4. describe the procedures used to confirm that an outbreak has been successfully controlled/eradicated, including any restrictions on restocking;

7.3.5. give details of any compensation payments made available to farmers etc when animals are slaughtered for disease control/eradication purposes.

8. **Compliance with the Terrestrial Code**

In addition to the documentary evidence that the provisions of appendix 3.8.3 are properly implemented and supervised, the Delegate of the country must submit a declaration indicating that in the zone:

8.1. no clinical CBPP has been detected for at least 2 years;

8.2. no CBPP vaccines have been used for at least 2 years in any susceptible species;

8.3. the country operates both clinical surveillance and disease reporting systems for CBPP adequate to detect clinical disease if it were present in the zone;

8.4. all clinical and pathological suggestive of CBPP is investigated by field and laboratory methods (including serological assessment) to refute a possible diagnosis of CBPP;

8.5. there are effective measures in force to prevent the re-introduction of the disease.

9. **Recovery of status**

Countries applying for recovery of status should comply with the provisions of Article 2.3.15.3 of the Terrestrial Code and provide detailed information as specified in sections 3.1, 3.2, 3.3, 5.2, 5.3 and 5.4 of this report. Information in relation to other sections need only be supplied if relevant.