The OIE Terrestrial Animal Health Standards Commission (hereafter referred to as the Terrestrial Code Commission) met at the OIE Headquarters in Paris from 19 to 30 September 2005.

The members of the Terrestrial Code Commission are listed in Appendix I. The agenda adopted is given in Appendix II.

The Director General of the OIE, Dr B. Vallat, welcomed the members and thanked them all for their willingness to participate in this important OIE work. He recalled the significant changes to the meeting timetable of the two Code Commissions in that the OIE was reverting to a two-year cycle for the preparation and adoption of standards, except in the case of international crises. He hoped that the changed timetable would further improve coordination in standards work between the Terrestrial Code Commission and the other Commissions.

Dr Vallat recalled the commitments made to Member Countries at the 73rd General Session regarding progress on some important texts which had been adopted on the understanding that outstanding Member Countries’ comments would be addressed. This included surveillance for bovine spongiform encephalopathy (BSE) (which would allow the official recognition of BSE status under the new chapter), bluetongue surveillance, the newly adopted standards on animal welfare, the ‘under study’ parts of the avian influenza chapter, and compartmentalisation. Dr Vallat also noted the need to address the recommendations arising from Regional Commission and other OIE meetings, including suggestions for improving the OIE Terrestrial Animal Health Code (hereafter referred to as the Terrestrial Code) chapters on Veterinary Services (including statutory body responsibilities, rapid response capability and auditing mechanisms).

Dr Vallat encouraged the Terrestrial Code Commission and the Aquatic Animal Health Standards Commission (hereafter referred to as the Aquatic Animals Commission) to continue their collaborative work on harmonisation of the two Codes.

The President of the Terrestrial Code Commission considered that the meeting was an appropriate time to examine the ways in which the Terrestrial Code Commission was operating and how procedures could be improved, particularly in relation to the other Commissions and the revised meeting schedule.

The Terrestrial Code Commission examined various Terrestrial Code texts in the light of Member Countries’ comments received just prior to and following the 73rd General Session. The outcome of the Terrestrial Code Commission’s work is presented as appendices to this report. Amendments made to existing chapters and previously circulated drafts are shown as double underlined text, with deleted text in strikeout.
The Terrestrial Code Commission thanked the following Member Countries for providing comments: Argentina, Australia, Botswana, Brazil, Canada, Chile, the European Union (EU), Japan, New Zealand, the Southern Cone countries of South America, Sudan, Switzerland, Taipei China, Thailand and the United States of America (USA).

The Terrestrial Code Commission strongly encouraged Member Countries to participate in the development of the OIE’s international standards by sending comments on this report in sufficient time for them to be considered by the Commission. It would assist the Terrestrial Code Commission if comments were submitted as specific proposed text changes, supported by a scientific rationale.

Comments need to reach the OIE Headquarters by 17 February 2006 in order to be considered at the next meeting of the Commission in March 2006. However, in order to meet the deadlines for meetings of the Animal Production Food Safety Working Group and the Scientific Commission on Animal Diseases (hereafter referred to as the Scientific Commission) comments on Appendices VII, XII, XXIV and XXV should reach the OIE Headquarters by 3 January 2006.

Member Countries should note that, unless stated otherwise, all texts submitted for comment in this report (Part A) may be proposed for adoption at the 74th General Session. Depending on the nature of the comments received on each text, the Terrestrial Code Commission will indicate in its March 2006 meeting report whether a particular text will be proposed for adoption or held over for further work.

A. TEXTS WHICH ARE SUBMITTED FOR MEMBER COUNTRY COMMENT

1. General definitions (Chapter 1.1.1.)

After examining a comment from Australia on the definition of ‘Case’, the Terrestrial Code Commission reconfirmed its previous position that limiting the pathogen by referring to ‘listed by the OIE’ would not be wise in order to encourage reporting of diseases not listed by the OIE, notably emerging diseases.

The suggestion by the EU to modify ‘Outbreak of disease or infection’ to be serially numbered was not adopted as the Terrestrial Code Commission believes that every Member Country has its own system of numbering and any numbering system OIE suggests may cause confusion.

After considering the comment by Portugal during the 73rd General Session, the definition of ‘Quarantine station’ was modified to accommodate disease specific conditions.

A set of definitions proposed by the Working Group of Animal Welfare was reviewed and endorsed by the Terrestrial Code Commission with some minor modifications.

Suggested changes are at Appendix III for the comment of Member Countries.

2. Evaluation of Veterinary Services (Chapters 1.3.3. and 1.3.4.)

The Terrestrial Code Commission received from the President of the ad hoc Group a draft revised text of Chapters 1.3.3. and 1.3.4. on the evaluation of Veterinary Services. This revision included recommendations on the evaluation of the Veterinary Statutory Body and on a procedure whereby a Member Country can request the OIE to organise an evaluation of its Veterinary Services.

The Terrestrial Code Commission also reviewed the tool (Performance, Vision, Strategy [PVS] Instrument) developed by the OIE and the Inter-American Institute for Cooperation on Agriculture (IICA) for the evaluation of Veterinary Services. This evaluation tool has been tested in the Americas and later reviewed and updated on the basis of this experience for a broader application. The Instrument was designed to indicate the areas of strength and weakness of a Veterinary Service (with a view to the allocation of resources) rather than to pass or fail it. The Terrestrial Code Commission was of the opinion that this tool could be used as a guide for self-evaluation by a Member Country of its Veterinary Services and for evaluation by the OIE of a Member Country’s Veterinary Services on a voluntary basis.

OIE Terrestrial Animal Health Standards Commission/September 2005
Suggested changes to the chapters are at Appendices IV and V for the comment of Member Countries. The current version of the PVS Instrument is attached for the comment of Member Countries (Appendix VI).

3. **Zoning and compartmentalisation (Chapter 1.3.5.)**

Comments received from Member Countries as well as issues raised during the 73rd General Session were examined.

Paragraphs in the articles on ‘Introduction’ and ‘General considerations’ were reorganised to address the issues more logically. The Terrestrial Code Commission also clarified the commitment of the Veterinary Administration by modifying the last paragraph of Article 1.3.5.2.

The Terrestrial Code Commission was concerned that the concept of compartmentalisation was not yet well understood. It noted that the application of compartmentalisation was not mandatory and it should be used in a similar manner to zoning, depending on the epidemiology of the disease. While the primary criteria for zoning are related to geography, those for compartments relate to biosecurity management measures; however, the application of zoning includes some biosecurity elements and compartmentalisation will involve a spatial element for some diseases.

The Terrestrial Code Commission also discussed some issues on compartmentalisation with the ad hoc Group on Epidemiology which is preparing an explanatory paper on the concept in order to provide guidance to Member Countries.

The revised Chapter 1.3.5. is submitted at Appendix VII for the comment of Member Countries.

4. **General guidelines for animal health surveillance (Appendix 3.8.1.)**

The Terrestrial Code Commission was advised that the Scientific Commission will review the work of the ad hoc Group on Epidemiology at the Commission’s meeting in January 2006 and present a revised appendix for consideration at the Terrestrial Code Commission’s meeting in March 2006. The Terrestrial Code Commission has requested the Scientific Commission to take into account comments received from Member Countries.

The Terrestrial Code Commission has also requested that some guidelines on surveillance for vectors be included in the appendix.

5. **Criteria for listing diseases (Chapter 2.1.1.)**

The Terrestrial Code Commission met with Dr K. Ben Jebara, Head of the OIE Animal Health Information Department, to discuss the comments from Member Countries on the criteria for listing diseases. Minor changes were made to Article 2.1.1.1. and the decision tree in Article 2.1.1.2. was amended accordingly. Member Countries’ proposals for the inclusion or deletion from the OIE list of diseases will be considered by the ad hoc Group on Animal Disease Notification. Member Countries are reminded that they need to submit a supporting statement (addressing the relevant criterion) with each proposal.

The revised text at Appendix VIII is submitted for the comment of Member Countries.

6. **Foot and mouth disease (Chapter 2.2.10. and Appendix 3.8.7.)**

The following modifications, in addition to some minor changes, were made to the chapter on foot and mouth disease (FMD) in response to Member Countries’ comments.

After examining a comment from the EU, the Terrestrial Code Commission modified the conditions for “Recovery of free status” in Article 2.2.10.7. to clarify that it also applies to country, not only to zone. After examining a comment from New Zealand, the Terrestrial Code Commission clarified point 5) of Article 2.2.10.8. The Terrestrial Code Commission discussed the comment from the EU regarding the need to certify the vaccination status of an animal and decided to ask the Scientific Commission to further examine the need for such a requirement in Articles 2.2.10.9. and 2.2.10.10.
The Terrestrial Code Commission decided to send the comments from the EU on the milk and milk products for animal feeding in Article 2.2.10.24. and on skins and trophies from wild susceptible animals in Article 2.2.10.29. to the Scientific Commission, for further examination.

The Terrestrial Code Commission noted that the revised Appendix 3.8.7. prepared by the Scientific Commission did not include the concept of compartmentalisation. As a result, the Terrestrial Code Commission did not incorporate the concept into the chapter as requested by some Member Countries.

Some surveillance issues raised by Member Countries were referred to the Scientific Commission for consideration.

The revised chapter and appendix are presented at Appendices IX and X for the comment of Member Countries.

7. **Bluetongue (Chapter 2.2.13.)**

After reviewing comments from New Zealand, the Terrestrial Code Commission modified the southern latitude boundary in Articles 2.2.13.1. and 2.2.13.2.

After examining comments from the EU on the distance from the infection front in which surveillance was required, the Terrestrial Code Commission modified the paragraph to give more flexibility, with a linkage to the proposed surveillance appendix on bluetongue.

The newly developed surveillance appendix was received from the Scientific Commission and is presented unchanged to Member Countries.

The revised chapter and appendix are presented at Appendices XI and XII for the comment of Member Countries.

8. **Bovine tuberculosis (Chapter 2.3.3.)**

The Terrestrial Code Commission reviewed comments from the EU and New Zealand. The Terrestrial Code Commission decided to forward all comments to the Scientific Commission for examination, including the proposal from some Member Countries to expand the scope of this chapter or develop a new chapter to include bison, deer and wildlife.

9. **Bovine spongiform encephalopathy (Chapter 2.3.13. and Appendix 3.8.4.)**

    a) **Chapter 2.3.13.**

    The Terrestrial Code Commission recalled the discussion at the 73rd General Session where some Member Countries were opposed to inclusion of muscle meat and blood products in the list of commodities which can be traded safely. However, arguments were largely based on studies using laboratory strains of transmissible spongiform encephalopathy (TSE) in laboratory animals, and many scientific papers have confirmed that different TSEs behave differently in various animal models. With respect to BSE, cattle provide the appropriate model to study the distribution of the agent in cattle. A number of studies has failed to demonstrate the presence of BSE in muscle meat or in the blood of experimentally infected cattle not showing clinical signs of BSE.

    The Terrestrial Code Commission also took into account information arising from a recent World Health Organization (WHO) Consultation on Tissue Infectivity Distribution in Transmissible Spongiform Encephalopathies, but noted that, in many cases, this was preliminary information requiring further validation. The meeting also referred to an equivocal result for muscle meat (semitendinosus muscle) arising from a clinically affected cow. The Terrestrial Code Commission did not believe that any changes in this regard were required to the current text, but it would continue to monitor the progress in the research.
Comments from New Zealand and the Southern Cone countries of South America, which were endorsed by the ad hoc Group on Surveillance for BSE, were taken into account and references to other TSEs was deleted from the BSE chapter, because there is little evidence that surveillance information on other TSEs is necessary to determine the risks presented by the BSE agent. Any risks presented by other TSEs are addressed by the application of measures such as feed bans.

The Terrestrial Code Commission has been concerned for some time that up-to-date recommendations needed to be developed for other commodities, and will ask the Director General to convene an expert group to examine the safety of gelatine and tallow.

It also considered the issue of the specifications in point g) of Article 2.3.13.1 and “…30 months of age or less…” was removed because there is no scientific basis for this age restriction. The reference to “…and were not suspect or confirmed BSE cases…” was also deleted as the ante-mortem and post-mortem inspection specified in the same sentence would automatically exclude such animals. Likewise, in Article 2.3.13.11, point 1)(a) was deleted, as point 1)(c) precludes any suspect or confirmed BSE cases.

The Terrestrial Code Commission did not agree with the EU comment that the requirement for annual review of the risk assessment in Article 2.3.13.2.was onerous. It noted that the requirement was that documentation be provided to indicate whether the situation had changed in the previous 12 months.

In revising Articles 2.3.13.3. and 2.3.13.4., the Terrestrial Code Commission took into consideration comments from Canada, the EU, New Zealand and Switzerland, as well as recommendations made by the ad hoc Group on Surveillance for BSE. Point 2) of Article 2.3.13.4. was modified to incorporate references to Type B surveillance. In response to a submission from New Zealand and after considering advice from an expert, the reference to progeny was deleted from point 3)(b)(iii) of Articles 2.3.13.3. and 2.3.13.4. and point 2 a) of Article 2.3.13.8.

The Terrestrial Code Commission is working with an expert to update the existing supporting document on BSE.

The revised chapter is at Appendix XIII for the comment of Member Countries.

b) Appendix 3.8.4.

The report of the second meeting of the ad hoc Group on Surveillance for BSE is at Appendix XXXIII for the information of Member Countries.

The Terrestrial Code Commission examined the appendix proposed by the experts and endorsed it with minor changes.

The Appendix on surveillance for BSE is at Appendix XIV for the comment of Member Countries.

c) Appendix 3.8.5.

The Terrestrial Code Commission recognised that, as a result of changes made in the Appendix 3.8.4., Appendix 3.8.5. needs to be revised. It will work on this and present the draft as a part of the report of its March 2006 meeting.

10. Classical swine fever (Chapter 2.6.7. and Appendix 3.8.8.)

At the request of several Member Countries, the Terrestrial Code Commission worked on the chapter on classical swine fever (CSF) to incorporate the concept of compartmentalisation. The chapter was modified in order to better harmonise various articles, including with equivalent articles in the FMD chapter. However, new science was not introduced.
The chapter now does not make specific reference to countries or zones where there is a different health status of the domestic and wild pig populations, unless compartmentalisation is applied to maintain separation of domestic from wild pigs.

Requests from New Zealand and Japan to refer to surveillance in point 2)b) of Article 2.6.7.4. have been addressed as a result of the revision of the article.

The prescriptive text was deleted from Article 2.6.7.6., as it was considered inappropriate to prescribe such conditions which should be developed on a case-by-case basis.

The Terrestrial Code Commission is submitting the modified chapter for Member Countries’ comment (Appendix XV), and will also submit it to the Scientific Commission to allow it to make the necessary changes to the appendix on surveillance. Among the changes required in the appendix, particular emphasis will need to be placed on the type of surveillance necessary to support the establishment and maintenance of a free compartment within infected countries or zones.

The Terrestrial Code Commission is awaiting advice from the Scientific Commission regarding commodities which could be safely traded regardless of the CSF status of the exporting country.

11. Avian influenza (Chapter 2.7.12. and Appendix 3.8.9.)

a) Chapter 2.7.12.

During the 73rd General Session, a revised Terrestrial Code chapter on avian influenza was adopted by the OIE International Committee. This revised chapter and the comments received from Argentina, Australia, Chile, the EU, the International Egg Commission, Japan, New Zealand and an expert were considered by the Terrestrial Code Commission. Among a number of general comments, in particular, comments on compartmentalisation and vaccination were taken into account when addressing specific articles.

New Zealand proposed that a new first article be drafted stating that eggs and poultry meat for human consumption can be freely traded from flocks not free from low pathogenic avian influenza (LPAI). The Terrestrial Code Commission did not think such an article was possible at this stage; however, it expanded on recommendations within relevant articles.

Japan proposed that the chapter should distinguish between “NAI free with vaccination” and “NAI free without vaccination”. Comments from Argentina and Chile appeared to support the Japanese proposal. The Terrestrial Code Commission noted that Appendix 3.8.9. addressed the issue and inserted reference to this appendix in the text.

New Zealand proposed that Article 2.7.12.6. be deleted. The Terrestrial Code Commission did not accept this proposal because live birds other than poultry pose an avian influenza risk to poultry.

b) Appendices

The Terrestrial Code Commission had a meeting with the ad hoc Group on Epidemiology to discuss their recommendations from their May 2005 meeting. The ad hoc Group confirmed that comments made by New Zealand had been considered, but had not resulted in changes to the text.

Following a comment submitted by an expert, the fifth paragraph of Article 3.8.9.7. was reorganised to better demonstrate how to distinguish vaccinated from infected poultry.

The Terrestrial Code Commission made some modifications to point 2)a) of Article 3.8.9.2., point 1) of Article 3.8.9.3. and Article 3.8.9.5. for clarification or consistency of terminology.
The Terrestrial Code Commission drafted a new appendix on procedures for the inactivation of highly pathogenic notifiable avian influenza (HPNAI) virus. The information in the appendix was compiled from a published paper and from a manuscript in press, provided by an expert. There are:


A revised chapter and appendix, and the new appendix on virus inactivation, are presented (Appendices XVI, XVII and XVIII) for the comment of Member Countries.

12. Semen and embryo related matters (Appendix 3.2.1)

The Terrestrial Code Commission accepted the comment from the EU and clarified point 1) of Article 3.2.1.3.

After examining the comment from Australia, the Terrestrial Code Commission modified Articles 3.2.1.5. and 3.2.1.6. to clarify that testing is unnecessary for animals in free countries.

In response to the comments from Australia, the Terrestrial Code Commission agreed to delete caseous lymphadenitis and border disease from point 1) of Article 3.2.1.6., as such diseases are not considered transmissible by semen.

After examining the comment from Australia, point 3) of Article 3.2.1.10. was modified to give further security to semen stored for export.

The revised appendix is at Appendix XIX for the comment of Member Countries.

13. Small hive beetle of honey bees (*Aethina tumida*) (Section 2.9.)

The Terrestrial Code Commission recalled the request from some Member Countries that a new chapter on this beetle be developed. The Terrestrial Code Commission examined a draft text prepared by the EU and a risk assessment prepared by a New Zealand expert, and decided to ask the Scientific Commission to develop a chapter on the small hive beetle of honey bees for consultation with Member Countries.

14. Animal welfare (Section 3.7.)

The Terrestrial Code Commission examined and endorsed the work of the Working Group on Animal Welfare in revising the four adopted chapters on animal welfare, taking into account comments received from Member Countries prior to the 73rd General Session, and the discussion at the General Session. The report of the fourth meeting of the Working Group on Animal Welfare is at Appendix XXXV for the information of Member Countries.

The Terrestrial Code Commission noted that some technical issues raised with experts in the ad hoc groups had not yet been responded to, but it expected that these should be finalised in time for the next meeting of the Terrestrial Code Commission in March 2006.

The four revised chapters are at Appendices XX, XXI, XXII and XXIII for the comment of Member Countries.

15. Animal production food safety

The Terrestrial Code Commission examined the report of the March 2005 meeting of the Working Group on Animal Production Food Safety and decided to circulate it for the information of Member Countries (Appendix XXXVI).
The Terrestrial Code Commission noted the work of the Food and Agriculture Organization (FAO) on good agricultural practice and recommended that, with regard to the Working Group document ‘Guide to good farming practices’, the OIE and the FAO coordinate their work with the aim of the information being published by both organisations for the guidance of Member Countries and the public.

Using a detailed discussion paper developed by the Working Group, the Terrestrial Code Commission drafted guidelines for the control of hazards of animal health and public health importance through ante-and post-mortem meat inspection. It is the Commission’s intention that this text become a Code chapter in a new section on food safety. The draft is at Appendix XXIV for Member Countries’ comment.

The Terrestrial Code Commission examined briefly some draft international health certificates, developed under the Working Group’s work programme. The Terrestrial Code Commission noted that the Working Group would be examining these draft certificates at its next meeting at the end of January 2006.

16. Animal identification and traceability

The Terrestrial Code Commission noted the report of the ad hoc Group on Animal Identification and Traceability which is at Appendix XXXIV for the information of Member Countries. The Terrestrial Code Commission supported the recommendation of the ad hoc Group that the general principles form a part of a horizontal chapter on animal identification and traceability, in conjunction with more detailed articles on essential elements. The Terrestrial Code Commission noted the importance of coordination with the Codex Commission during the development of these standards.

The Terrestrial Code Commission made some modifications to the proposed definitions and general principles for animal identification and traceability, and is presenting the modified text at Appendix XXV for Member Countries’ comment.

17. Carcass disposal

The Terrestrial Code Commission received from the Scientific Commission for Animal Diseases a revised text on the disposal of carcasses. The original text had been circulated for Member Countries’ comment in 2004. The Terrestrial Code Commission proposed significant changes to the text (including to the title) and decided that the changes should be reviewed by the Scientific Commission before being circulated for further comment by Member Countries.

18. Paratuberculosis (Chapter 2.2.6.)

The Terrestrial Code Commission recalled the discussion on paratuberculosis at the 68th General Session in which Member Countries expressed their concern over possible trade implications of a proposed revised text. The text previously circulated for comment is at Appendix XXXVII for information.

As the current Terrestrial Code chapter is without technical content, the Terrestrial Code Commission is seeking advice from Member Countries on how to proceed in the development of an up-to-date chapter on paratuberculosis, with recommendations which do not unnecessarily disrupt trade.

19. Equine diseases (Section 2.5.)

After consultation with OIE Reference Laboratories on some equine disease chapters in need of updating, the Terrestrial Code Commission modified chapters on equine infectious anaemia, equine piroplasmosis, equine rhinopneumonitis, glanders and equine viral arteritis. The modified texts, at Appendices XXVI, XXVII, XXVIII, XXIX and XXX, are presented for comment by Member Countries.

The Terrestrial Code Commission examined a proposed revised chapter on African horse sickness prepared by a group of experts, and made some appropriate changes. The proposed revision (which is based on the bluetongue chapter), is circulated as clean text at Appendix XXXI for the comment of Member Countries. It will also be sent to the Scientific Commission for consideration.
20. **Bovine viral diarrhoea-mucosal disease**

The President of the Commission met some European experts on bovine viral diarrhoea-mucosal disease (BVD-MD) to discuss the BVD-MD situation, control efforts in Europe and possible future steps to make the European experience known internationally. As a result, the Terrestrial Code Commission discussed how guidance on the disease could be provided to Member Countries. It noted that a chapter had not been developed, but that some recommendations were present in the chapter on bovine and small ruminant semen.

The Terrestrial Code Commission recognised that BVD-MD has a worldwide distribution, but that, under certain circumstances, it can cause economic loss. It also recognised that, if certain procedures are followed, the disease could be controlled and eventually eradicated at a herd or regional level. The Terrestrial Code Commission concluded that it would not be appropriate to develop a specific chapter, but that guidance on control could be offered to Member Countries.

The Terrestrial Code Commission is therefore seeking advice from Member Countries as to how the OIE could address diseases such as BVD-MD and listeriosis, using alternative mechanisms to those currently used in the *Terrestrial Code* or *Manual* for such diseases, in order to provide Member Countries with useful guidance on managing such diseases without causing unjustified disruptions to trade.

21. **International transfer of pathogens (Chapter 1.4.5.)**

In consultation with the Terrestrial Code Commission, the Biological Standards Commission (hereafter referred to as the Laboratories Commission) decided to update Chapter 1.4.5. of the *Terrestrial Code* and the relevant sections of the *Terrestrial Manual*. The Terrestrial Code Commission expects to review revised text at its next meeting.

22. **Future work programme**

The Terrestrial Code Commission reviewed its work programme, taking into account the outcomes of the 73rd General Session, submissions received from Member Countries, and input from the Scientific Commission and the Laboratories Commission. A table summarising planned future activities for the Terrestrial Code Commission is at Appendix XXXII for the comment of Member Countries.

**B. REPORTS OF WORKING GROUPS AND AD HOC GROUPS**

The following reports are for the information of Member Countries:

- *Ad hoc Group on Surveillance for Bovine Spongiform Encephalopathy* ([Appendix XXXIII](#))
- *Ad hoc Group on Animal Identification and Traceability* ([Appendix XXXIV](#))
- *Animal Welfare Working Group* ([Appendix XXXV](#))
- *Animal Production Food Safety Working Group* ([Appendix XXXVI](#))

**C. OTHER DOCUMENT**

The following document is for the information of Member Countries: Chapter 2.2.6. on paratuberculosis proposed in 2000 ([Appendix XXXVII](#)).

The list of chapters and appendices circulated for the comment of Member Countries is in Section A of this report.

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*OIE Terrestrial Animal Health Standards Commission/September 2005*
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MEETING OF THE OIE TERRESTRIAL ANIMAL HEALTH STANDARDS COMMISSION

Paris, 19-30 September 2005

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Agenda

Item 1  General definitions (Chapter 1.1.1.)
Item 2  Evaluation of Veterinary Services (Chapters 1.3.3. and 1.3.4.)
Item 3  Zoning and compartmentalisation (Chapter 1.3.5.)
Item 4  General guidelines for animal health surveillance (Appendix 3.8.1.)
Item 5  Criteria for listing diseases (Chapter 2.1.1.)
Item 6  Foot and mouth disease (Chapter 2.2.10. and Appendix 3.8.7.)
Item 7  Bluetongue (Chapter 2.2.13. and surveillance appendix)
Item 8  Bovine tuberculosis (Chapter 2.3.3.)
Item 9  Bovine spongiform encephalopathy (Chapter 2.3.13. and Appendix 3.8.4.)
Item 10  Classical swine fever (Chapter 2.6.7. and Appendix 3.8.8.)
Item 11  Avian influenza (Chapter 2.7.12. and Appendix 3.8.9.)
Item 12  Semen and embryo related matters (Appendix 3.3.5. and Appendix 3.2.1.)
Item 13  Small hive beetle on honey bees (Section 2.9.)
Item 14  Animal welfare (Section 3.7)
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Item 16  Animal identification and traceability
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Item 18  Paratuberculosis (Chapter 2.2.6.)
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Item 20  Bovine viral diarrhoea-mucosal disease
Item 21  International transfer of pathogens (Chapter 1.4.5.)
Item 22  Work programme
Item 23  Others
CHAPTER 1.1.1.

GENERAL DEFINITIONS

Animal handler
A person with a knowledge of the behaviour and needs of animals which, with appropriate experience and a professional and positive response to an animal’s needs, results in effective management and good welfare. Their competence should be demonstrated through independent assessment and certification.

Container
A non-self-propelled receptacle or other rigid structure for holding animals during a journey by one or several means of transport.

Death
Irreversible loss of brain activity demonstrable by the loss of brain stem reflexes.

Journey
An animal transport journey commences when the first animal is loaded onto a vehicle/vessel or into a container and ends when the last animal is unloaded, and includes any stationary resting / holding periods of less than 48 hours. The same animals do not commence a new journey until after a period of over 48 hours for rest and recuperation, with adequate feed and water.

Killing
Any procedure which causes the death of an animal.

Lairage
Pens, yards and other holding areas used for accommodating animals in order to give them necessary attention (including water, feed, rest) before they are moved on or used for specific purposes including slaughter.

Loading/Unloading
Loading: the procedure of moving animals onto a vehicle/vessel or into a container for transport purposes; unloading: the procedure of moving animals off a vehicle/vessel or out of a container.

Post-journey period
The period between unloading and either recovery from the effects of the journey or slaughter (if this occurs before recovery).

Pre-journey period
The period during which animals are identified, and often assembled for the purpose of loading them.

Resting point
A place where the journey is interrupted to rest, feed or water the animals; the animals may remain in the vehicle/vessel or container, or be unloaded.

Restraint
The application to an animal of any procedure designed to restrict its movements.

Slaughter
Any procedure which causes the death of an animal by bleeding.

Space allowance
The measure of the floor area and height on a vehicle/vessel or container allocated per individual or body weight of animals transported.
Appendix III (contd)

**Stocking density**

The number or body weight of animals per unit area on a *vehicle/vessel or container*.

**Stunning**

Any mechanical, electrical, chemical or other procedure which causes immediate loss of consciousness; when used before slaughter, the loss of consciousness lasts until death from the slaughter process; in the absence of slaughter, the procedure would allow the animal to recover consciousness.

**Transport**

The procedures associated with the carrying of animals for commercial purposes from one location to another by land (road and rail), sea or air.

**Transporter**

The person licensed by the *Competent Authority to transport animals*.

**Travel**

The movement of a *vehicle/vessel or container* carrying animals from one location to another.

**Vehicle/vessel**

Any means of conveyance including train, truck, *aircraft* or ship that is used for carrying animal(s).

**Slaughterhouse/abattoir**

Premises, including facilities for moving or lairaging animals, used for the slaughter of animals for human consumption or animal feeding, and approved by the *Veterinary Services* or other *Competent Authority*.

**Quarantine station**

A facility under the control of the *Veterinary Authority* where a group of animals is maintained in isolation with no direct or indirect contact with other animals, to prevent the transmission of specified disease(s), in order to while the animals are undergoing observation for a specified length of time and, if appropriate, testing and treatment.
CHAPTER 1.3.3.

EVALUATION OF VETERINARY SERVICES

Article 1.3.3.1.

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

Compliance with these fundamental principles by the Veterinary Services of a Member Country is important to the establishment and maintenance of confidence in its international veterinary certificates by the Veterinary Services of other Member Countries.

The same fundamental principles should apply in countries where the responsibility for establishing or applying certain animal health measures, or issuing some international veterinary certificates is exercised by an organisation other than the Veterinary Services, or by an authority or agency on behalf of the Veterinary Services. In all cases, the Veterinary Services retain ultimate responsibility for the application of these principles.

These fundamental principles are presented in Article 1.3.3.2. The remaining Other factors affecting quality are described in Part 1. (notification, principles of certification, etc.). and the document entitled Guidelines for the evaluation of Veterinary Services included in Chapter 1.3.4.

The quality of Veterinary Services can be measured through an evaluation, whose general principles are described in Article 1.3.3.3. and in Article 1.3.3.4.

Guidelines for the evaluation of Veterinary Services are described in Chapter 1.3.4.

A procedure for evaluating Veterinary Services by OIE experts, on a voluntary basis, is described in Article 1.3.3.5.

Article 1.3.3.2.

Fundamental principles of quality

The Veterinary Services shall comply with the following principles to ensure the quality of their activities:

1. **Professional judgement**

   The personnel of Veterinary Services should have the relevant qualifications, scientific expertise and experience to give them the competence to make sound professional judgements.

2. **Independence**

   Care should be taken to ensure that Veterinary Services personnel are free from any commercial, financial, hierarchical, political or other pressures which might affect their judgement or decisions.
Appendix IV (contd)

3. Impartiality

The Veterinary Services should be impartial. In particular, all the parties affected by their activities have a right to expect their services to be delivered under reasonable and non-discriminatory conditions.

4. Integrity

The Veterinary Services should guarantee that the work of each of their personnel is of a consistently high level of integrity. Any fraud, corruption or falsification should be identified and corrected.

5. Objectivity

The Veterinary Services should at all times act in an objective, transparent and non-discriminatory manner.

6. General organisation

The Veterinary Services must be able to demonstrate by means of appropriate legislation, sufficient financial resources and effective organisation that they are in a position to have control of the establishment and application of animal health measures, and of international veterinary certification activities. Legislation should be suitably flexible to allow for judgements of equivalence and efficient responses to changing situations. In particular, they should define and document the responsibilities and structure of the organisations in charge of the animal identification system, control of animal movements, animal disease control and reporting systems, epidemiological surveillance and communication of epidemiological information.

A similar demonstration should be made by Veterinary Services when they are in charge of veterinary public health activities.

The Veterinary Services should have at their disposal effective systems for animal disease surveillance and for notification of disease problems wherever they occur, in accordance with the provisions of this Terrestrial Code. Adequate coverage of animal populations should also be demonstrated. They should at all times endeavour to improve their performance in terms of animal health information systems and animal disease control.

The Veterinary Services should define and document the responsibilities and structure of the organisation (in particular the chain of command) in charge of issuing international veterinary certificates.

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

7. Quality policy

The Veterinary Services should define and document their policy and objectives for, and commitment to, quality, and should ensure that this policy is understood, implemented and maintained at all levels in the organisation. Where conditions allow, they may implement a quality system corresponding to their areas of activity and appropriate for the type, range and volume of work that they have to perform. The guidelines for the quality and evaluation of Veterinary Services propose a suitable reference system, which should be used if a Member Country choose to adopt a quality system.
8. Procedures and standards

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

a) programming and management of activities, including international veterinary certification activities;
b) prevention, control and notification of disease outbreaks;
c) risk analysis, epidemiological surveillance and zoning;
d) inspection and sampling techniques;
e) diagnostic tests for animal diseases;
f) preparation, production, registration and control of biological products for use in the diagnosis or prevention of diseases;
g) border controls and import regulations;
h) disinfection and disinfestation;
i) treatments intended to destroy, if appropriate, pathogens in animal products.

Inasmuch as the OIE has adopted standards on these matters, the Veterinary Services should comply with these standards when applying animal health measures and when issuing international veterinary certificates.

9. Information, complaints and appeals

The Veterinary Administration should undertake to reply to legitimate requests from Veterinary Administrations of other Member Countries or any other authority, in particular ensuring that any requests for information, complaints or appeals that they may present are dealt with in a timely manner.

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

10. Documentation

The Veterinary Services should have at their disposal a reliable and up to date documentation system suited to their activities.

11. Self-evaluation

The Veterinary Services should undertake periodical self-evaluation especially by documenting achievements against goals, and demonstrating the efficiency of their organisational components and resource adequacy.
Appendix IV (contd)

A Member Country can request the Director General of the OIE to arrange for an expert or experts to assist in the process.

A procedure for evaluating Veterinary Services by OIE experts, on a voluntary basis, is described in Article 1.3.3.5.

12. Communication

Veterinary Services should have effective internal and external systems of communication covering administrative and technical staff and parties affected by their activities.

13. Human and financial resources

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

Article 1.3.3.3.

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

Any evaluation of Veterinary Services should be conducted having regard to the OIE Guidelines for the evaluation of Veterinary Services presented in Chapter 1.3.4. of this Terrestrial Code.

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

Article 1.3.3.4.

A Member Country which intends to conduct an evaluation of another Member Country's Veterinary Services should give them notice in writing. This notice should define the purpose of the evaluation and details of the information required.

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

The evaluation process should take into account the fundamental principles and other factors of quality laid down in Article 1.3.3.1. and in Article 1.3.3.2. It should also take into consideration the specific circumstances regarding quality, as described in Article 1.3.3.1., prevailing in the countries concerned.
Appendix IV (contd)

The outcome of the evaluation conducted by a Member Country should be provided in writing as soon as possible, and in any case within 4 months of receipt of the relevant information, to the Member Country which has undergone the evaluation. The evaluation report should detail any findings which affect trade prospects. The Member Country which conducts the evaluation should clarify in detail any points of the evaluation on request.

In the event of a dispute between two Member Countries over the conduct or the conclusions of the evaluation of the Veterinary Services, the matter should be dealt with having regard to the procedures set out in Article 1.3.1.3.

Article 1.3.3.5.

Voluntary evaluation by OIE experts

The OIE maintains a procedure for the evaluation of the Veterinary Services of a Member Country, on a voluntary basis.

Under this procedure, on the receipt of a request from a Member Country, the Director General of the OIE recommends an expert(s) from a list of evaluators approved by the OIE International Committee.

The expert(s) evaluates the Veterinary Services of the Member Country against the provisions in Chapter 1.3.4 of the Terrestrial Code, using the Performance, Vision, Strategy [PVS] Instrument as a guide, and produces a report.

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

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CHAPTER 1.3.4.

GUIDELINES FOR THE EVALUATION OF VETERINARY SERVICES

Article 1.3.4.1.

General considerations

1. Evaluation of Veterinary Services is an important element in the risk analysis process which countries may legitimately use in their policy formulations directly applying to animal health and sanitary controls of international trade in animals, animal-derived products, animal genetic material and animal feedstuffs.

Any evaluation should be carried out with due regard for Chapter 1.3.3. of this Terrestrial Code.

2. In order to ensure that objectivity is maximised in the evaluation process, it is essential for some standards of discipline to be applied. The OIE has developed these guidelines which can be practically applied to the evaluation of Veterinary Services. These are relevant for evaluation of the Veterinary Services of one country by those of another country for the purposes of risk analysis in international trade. These guidelines (in conjunction with the Performance, Vision, Strategy [PVS] Instrument) will be used by OIE experts when conducting an evaluation on the request of a Member Country. The guidelines are also applicable for evaluation by a country of its own Veterinary Services – the process known as self-evaluation or self-assessment – and for periodic re-evaluation.

In carrying out a risk analysis prior to deciding the sanitary/zoosanitary conditions for the importation of a commodity, an importing country is justified in regarding its evaluation of the Veterinary Services of the exporting country as critical.

3. The purpose of evaluation may be either to assist a national authority in the decision-making process regarding priorities to be given to its own Veterinary Services (self-evaluation) or to assist the process of risk analysis in international trade in animals and animal-derived products to which official sanitary and/or zoosanitary controls apply.

4. In both situations, the evaluation should demonstrate that the Veterinary Services have the capability for effective control of the sanitary and zoosanitary status of animals and animal products. Key elements to be covered in this process include resource adequacy, management capability, legislative and administrative infrastructures, independence in the exercise of official functions and performance history, including disease reporting.

5. Competence and integrity are qualities on which others base their confidence in individuals or organisations. Mutual confidence between relevant official Veterinary Services of trading partner countries contributes fundamentally to stability in international trade in animals and animal-related products. In this situation, scrutiny is directed more at the exporting country than at the importing country.
Appendix V (contd)

6. Although quantitative data can be provided on Veterinary Services, the ultimate evaluation will be essentially qualitative. While it is appropriate to evaluate resources and infrastructure (organisational, administrative and legislative), it is also appropriate to place emphasis on the evaluation of the quality of outputs and performance of Veterinary Services. Evaluation should take into consideration any quality systems used by Veterinary Services.

7. An importing country has a right of assurance that information on sanitary/zoosanitary situations provided by the Veterinary Services of an exporting country is objective, meaningful and correct. Furthermore, the Veterinary Services of the importing country are entitled to expect validity in the veterinary certification of export.

8. An exporting country is entitled to expect that its animals and animal products will receive reasonable and valid treatment when they are subjected to import inspection in the country of destination. The country should also be able to expect that any evaluation of its standards and performance will be conducted on a non-discriminatory basis. The importing country should be prepared and able to defend any position which it takes as a consequence of the evaluation.

9. As the Veterinary statutory body is not a part of the Veterinary Services, an evaluation of that body should be carried out to ensure that the registration/licensing of veterinarians and authorisation of veterinary para-professionals is included.

Article 1.3.4.2.

Scope

1. In the evaluation of Veterinary Services, the following items may be considered, depending on the purpose of the evaluation:
   - organisation, structure and authority of the Veterinary Services;
   - human resources;
   - material (including financial) resources;
   - functional capabilities and legislative support;
   - animal health and veterinary public health controls;
   - formal quality systems including quality policy;
   - performance assessment and audit programmes;
   - participation in OIE activities and compliance with OIE Member Countries’ obligations.

2. To complement the evaluation of Veterinary Services, it is necessary to also consider the organisational structure and functioning of the Veterinary statutory body should also be considered.
3. Article 1.3.4.14. outlines appropriate information requirements for:

- self-evaluation by national *Veterinary Services* which perceive a need to prepare information for national or international purposes;

- evaluation by a prospective or actual *importing country* of the *Veterinary Services* of a prospective or actual *exporting country*;

- verification or re-verification of an evaluation in the course of a visit to the *exporting country* by the *importing country*.

4. The PVS Instrument should be used as a guide in conducting evaluations and self-evaluations.

**Article 1.3.4.3.**

**Evaluation criteria for the organisational structure of the Veterinary Services**

1. A key element in the evaluation is the study of the organisation and structure of the official *Veterinary Services*. The *Veterinary Services* should define and set out their policy, objectives and commitment to quality systems and standards. These organisational and policy statements should be described in detail. Organisational charts and details of functional responsibilities of staff should be available for evaluation. The role and responsibility of the Chief Veterinary Officer/Veterinary Director should be clearly defined. Lines of command should also be described.

2. The organisational structure should also clearly set out the interface relationships of government Ministers and departmental Authorities with the Chief Veterinary Officer/Veterinary Director and the *Veterinary Services*. Formal relationships with statutory authorities and with industry organisations and associations should also be described. It is recognised that Services may be subject to changes in structure from time to time. Major changes should be notified to trading partners so that the effects of re-structuring may be assessed.

3. Organisational components of *Veterinary Services* which have responsibility for key functional capabilities should be identified. These capabilities include epidemiological surveillance, disease control, import controls, animal disease reporting systems, animal identification systems, traceability systems, animal movement control systems, communication of epidemiological information, training, inspection and certification. Laboratory and field systems and their organisational relationships should be described.

4. To reinforce the reliability and credibility of their services, the *Veterinary Services* may have set up quality systems that correspond with their fields of activity and to the nature and scale of activities that they carry out. Evaluation of such systems should be as objective as possible.

5. The *Veterinary Administration* alone speaks for the country as far as official international dialogue is concerned. This is also particularly important to cases where zoning and regionalisation are being applied. The responsibilities of the national *Veterinary Administration* and all *Veterinary Authorities* in that country should be made clear in the process of evaluation of *Veterinary Services*. 
Appendix V (contd)

6. A Veterinary Authority is defined in Chapter 1.1.1. of this Terrestrial Code. As some countries have some official Veterinary Authority roles vested in autonomous sub-national (state/provincial, municipal) government bodies, there is an important need to assess the role and function of these Services. Details of their roles, relationship (legal and administrative) to each other and to the national Veterinary Services should be available for evaluation. Annual reports, review findings and access to other information pertinent to the animal health activities of such bodies should also be available.

7. Similarly, where the national Veterinary Services have arrangements with other providers of relevant services such as universities, laboratories, information services, etc., these arrangements should also be described. For the purposes of evaluation, it is appropriate to expect that the quality of organisational and functional standards which apply to Veterinary Services should also apply to the services of these other providers.

Article 1.3.4.4.

Evaluation criteria for quality systems

1. The Veterinary Services should demonstrate a commitment to the quality of the processes and outputs of their services. Where services or components of services are delivered under a formal quality systems programme which is based on OIE recommended standards or, especially in the case of laboratory components of Veterinary Services other internationally recognised quality standards, the Veterinary Services undergoing evaluation should make available evidence of accreditation, details of the documented quality processes and documented outcomes of all relevant audits undertaken.

2. Where the Veterinary Services undergoing evaluation make large use of formal quality systems in the delivery of their services, it is appropriate that greater emphasis be placed on the outcomes of evaluation of these quality systems than on the resource and infrastructural components of the services.

Article 1.3.4.5.

Evaluation criteria for human resources

1. The Veterinary Services should demonstrate that their human resource component includes an integral core of full-time civil service employees. This core must include veterinarians. It should also include administrative officials and veterinary para-professionals. The human resources may also include part-time and private sector veterinarians and veterinary para-professionals. It is essential that all the above categories of personnel be subject to legal disciplinary provisions. Data relating to the resource base of the Veterinary Services undergoing evaluation should be available.

2. In addition to raw quantitative data on this resource base, the functions of the various categories of personnel in the Veterinary Services should be described in detail. This is necessary for analysis and estimation of the appropriateness of the application of qualified skills to the tasks undertaken by the Veterinary Services and may be relevant, for example, to the roles of veterinarians and veterinary para-professionals in field services. In this case, the evaluation should provide assurances that disease monitoring is being conducted by a sufficient number of qualified, experienced field veterinarians who are directly involved in farm visits; there should not be an over-reliance on veterinary para-professionals for this task.
3. Analysis of these data can be used to estimate the potential of the *Veterinary Services* to have reliable knowledge of the state of animal health in the country and to support an optimal level of animal disease control programmes. A large population of private veterinarians would not provide the *Veterinary Services* with an effective epizootiological information base without legislative (e.g. compulsory reporting of notifiable diseases) and administrative (e.g. official animal health surveillance and reporting systems) mechanisms in place.

4. These data should be assessed in close conjunction with the other information described in this Chapter. For example, a large field staff (veterinarians and veterinary para-professionals) need fixed, mobile and budgetary resources for animal health activities in the livestock farming territory of the country. If deficiencies are evident, there would be reason to challenge the validity of epizootiological information.

**Article 1.3.4.6.**

**Evaluation criteria for material resources**

1. **Financial**

   Actual yearly budgetary information regarding the *Veterinary Services* should be available and should include the details set out in the model questionnaire outlined in Article 1.3.4.14. Information is required on conditions of service for veterinary staff (including salaries and incentives) and should provide a comparison with the private sector and perhaps with other professionals. Information should also be available on non-government sources of revenue available to veterinarians in their official responsibilities.

2. **Administrative**

   a) **Accommodation**

   The *Veterinary Services* should be accommodated in premises suitable for efficient performance of their functions. The component parts of the *Veterinary Services* should be located as closely as possible to each other at the central level, and in the regions where they are represented, in order to facilitate efficient internal communication and function.

   b) **Communications**

   The *Veterinary Services* should be able to demonstrate that they have reliable access to effective communications systems, especially for animal health surveillance and control programmes.

   Inadequate communications systems within the field services components of these programmes or between outlying offices and headquarters, or between the *Veterinary Services* and other relevant administrative and professional services, signify an inherent weakness in these programmes. Adequate communications systems between laboratories and between field and laboratory components of the *Veterinary Services* should also be demonstrated.
Examples of types of communications which should be routinely available on an adequate country-wide basis are national postal, freight and telephone networks. Rapid courier services, facsimile and electronic data interchange systems (e.g. e-mail and Internet services) are examples of useful communication services which, if available, can supplement or replace the others. A means for rapid international communication should be available to the national \textit{Veterinary Services}, to permit reporting of changes in national disease status consistent with OIE recommendations and to allow bilateral contact on urgent matters with counterpart \textit{Veterinary Services} in trading-partner countries.

c) Transport systems

The availability of sufficient reliable transport facilities is essential for the performance of many functions of \textit{Veterinary Services}. This applies particularly to the field services components of animal health activities (e.g. emergency response visits). Otherwise, the \textit{Veterinary Services} cannot assure counterpart services in other countries that they are in control of the animal health situation within the country.

Appropriate means of transport are also vital for the satisfactory receipt of samples to be tested at veterinary laboratories, for inspection of imports and exports, and for the performance of animals and animal product inspection in outlying production or processing establishments.

3. Technical

Details available on laboratories should include resources data, programmes under way as well as those recently completed and review reports on the role or functions of the laboratory. Information as described in the model questionnaire should be used in the evaluation of laboratory services.

a) Cold chain for laboratory samples and veterinary medicines

Adequate refrigeration and freezing systems should be available and should be used throughout the country to provide suitable low temperature protection for laboratory samples in transit or awaiting analysis, as well as veterinary medical products (e.g. vaccines) when these are required for use in animal disease control programmes. If these assurances cannot be given, it may be valid to discount many types of test results, as well as the effectiveness of certain disease control programmes and the export inspection system in the country undergoing evaluation.

b) Diagnostic laboratories

Analysis of the laboratory service component of \textit{Veterinary Services}, which would include official governmental laboratories and other laboratories accredited by the \textit{Veterinary Services} for specified purposes, is an essential element of the evaluation process. The quality of the veterinary diagnostic laboratories of a country underpins the whole control and certification processes of the zoosanitary/sanitary status of exported animals and animal products, and therefore these laboratories should be subject to rigid quality assurance procedures and should use international quality assurance programmes (wherever available) for standardising test methodologies and testing proficiency. An example is the use of International Standard Sera for standardising reagents.
This emphasis is valid whether one relates it to the actual testing performed on individual export consignments or to the more broad and ongoing testing regimes which are used to determine the animal health and veterinary public health profiles of the country and to support its disease control programmes. For the purposes of evaluation, veterinary diagnostic laboratories include those which are concerned with either animal health or veterinary public health activities. The *Veterinary Services* must approve and designate these laboratories for such purposes and have them audited regularly.

c) Research

The scope of animal disease and veterinary public health problems in the country concerned, the stages reached in the controls which address those problems and their relative importance can be measured to some degree by analysis of information on government priorities and programmes for research in animal health. This information should be accessible for evaluation purposes.

**Article 1.3.4.7.**

**Functional capabilities and legislative support**

1. **Animal health and veterinary public health**

   The *Veterinary Services* should be able to demonstrate that they have the capacity, supported by appropriate legislation, to exercise control over all animal health matters. These controls should include, where appropriate, compulsory notification of prescribed animal diseases, inspection, movement controls through systems which provide adequate traceability, registration of facilities, quarantine of infected premises/areas, testing, treatment, destruction of infected *animals* or contaminated materials, controls over the use of veterinary medicines, etc. The scope of the legislative controls should include domestic *animals* and their reproductive material, animal products, wildlife as it relates to the transmission of *diseases* to humans and domestic *animals*, and other products subject to veterinary inspection. Arrangements should exist for co-operation with the *Veterinary Authorities* of the neighbouring countries for the control of animal diseases in border areas and for establishing linkages to recognise and regulate transboundary activities. Information on the veterinary public health legislation covering the production of products of animal origin for national consumption may be also considered in the evaluation.

2. **Export/import inspection**

   National *Veterinary Services* should have appropriate legislation and adequate capabilities to prescribe the methods for control and to exercise systematic control over the import and export processes of *animals* and animal products in so far as this control relates to sanitary and *zoosanitary* matters. The evaluation should also involve the consideration of administrative instructions to ensure the enforcement of importing country requirements during the pre-export period.

   In the context of production for export of foodstuffs of animal origin, the *Veterinary Services* should demonstrate that comprehensive legislative provisions are available for the oversight by the relevant authorities of the hygienic process and to support official inspection systems of these *commodities* which function to standards consistent with or equivalent to relevant Codex Alimentarius and OIE standards.
Appendix V (contd)

Control systems should be in place which permit the exporting Veterinary Authorities to approve export premises. The Veterinary Services should also be able to conduct testing and treatment as well as to exercise controls over the movement, handling and storage of exports and to make inspections at any stage of the export process. The product scope of this export legislation should include, *inter alia*, animals and animal products (including animal semen, ova and embryos), and animal feedstuffs.

The national Veterinary Services should be able to demonstrate that they have adequate capabilities and legislative support for zoosanitary control of imports and transit of animals, animal products and other materials which may introduce animal diseases. This could be necessary to support claims by the Veterinary Services that the animal health status of the country is suitably stable, and that cross-contamination of exports from imports of unknown or less favourable zoosanitary status is unlikely. The same considerations should apply in respect of veterinary control of public health. The Veterinary Services should be able to demonstrate that there is no conflict of interest when certifying veterinarians are performing official duties.

Legislation should also provide the right to deny and/or withdraw official certification. Penalty provisions applying to malpractice on the part of certifying officials should be included.

The Veterinary Services should demonstrate that they are capable of providing accurate and valid certification for exports of animals and animal products, based on Section 1.2. of the Terrestrial Code. They should have appropriately organised procedures which ensure that sanitary/animal health certificates are issued by efficient and secure methods. The documentation control system should be able to correlate reliably the certification details with the relevant export consignments and with any inspections to which the consignments were subjected.

Security in the export certification process, including electronic documentation transfer, is important. A system of independent compliance review is desirable, to safeguard against fraud in certification by officials and by private individuals or corporations. The certifying veterinarian should have no conflict of interest in the commercial aspects of the animals or animal product being certified and be independent from the commercial parties.

**Article 1.3.4.8.**

**Animal health controls**

1. **Animal health status**

An updated assessment of the present animal disease status of a country is an important and necessary procedure. For this undertaking, studies of the OIE publications such as *World Animal Health*, the *Bulletin* and *Disease Information* must be fundamental reference points. The evaluation should consider the recent history of the compliance of the country with its obligations regarding international notification of animal diseases. In the case of an OIE Member Country, failure to provide the necessary animal health reports consistent with OIE requirements will detract from the overall outcome of the evaluation of the country.

An exporting country should be able to provide further, detailed elaboration of any elements of its animal disease status as reported to the OIE. This additional information will have particular importance in the case of animal diseases which are foreign to or strictly controlled in the importing country or region. The ability of the Veterinary Services to substantiate elements of their animal disease status reports with surveillance data, results of monitoring programmes and details of disease history is highly relevant to the evaluation. In the case of evaluation of the Veterinary Services of an exporting country for international trade purposes, an importing country should be able to demonstrate the reasonableness of its request and expectations in this process.
2. **Animal health control**

Details of current animal disease control programmes should be considered in the evaluation. These programmes would include epidemiological surveillance, official government-administered or officially-endorsed, industry-administered control or eradication programmes for specific diseases or disease complexes, and animal disease emergency preparedness. Details should include enabling legislation, programme plans for epidemiological surveillance and animal disease emergency responses, quarantine arrangements for infected and exposed animals or herds, compensation provisions for animal owners affected by disease control measures, training programmes, physical and other barriers between the free country or zone and those infected, incidence and prevalence data, resource commitments, interim results and programme review reports.

3. **National animal disease reporting systems**

The presence of a functional animal disease reporting system which covers all agricultural regions of the country and all veterinary administrative control areas should be demonstrated.

An acceptable variation would be the application of this principle to specific zones of the country. In this case also, the animal disease reporting system should cover each of these zones. Other factors should come to bear on this situation, e.g. the ability to satisfy trading partners that sound animal health controls exist to prevent the introduction of disease or export products from regions of lesser veterinary control.

**Article 1.3.4.9.**

**Veterinary public health controls**

1. **Food hygiene**

The national Veterinary Services should be able to demonstrate effective responsibility for the veterinary public health programmes relating to the production and processing of animal products. If the national Veterinary Services do not exercise responsibility over these programmes, the evaluation should include a comprehensive review of the role and relationship of the organisations (national, state/provincial, and municipal) which are involved. In such a case, the evaluation should consider whether the national Veterinary Services can provide guarantees of responsibility for an effective control of the sanitary status of animal products throughout the slaughter, processing, transport and storage periods.

2. **Zoonoses**

Within the structure of Veterinary Services, there should be appropriately qualified personnel whose responsibilities include the monitoring and control of zoonotic diseases and, where appropriate, liaison with medical authorities.

3. **Chemical residue testing programmes**

Adequacy of controls over chemical residues in exported animals, animal products and feedstuffs should be demonstrated. Statistically-based surveillance and monitoring programmes for environmental and other chemical contaminants in animals, in animal-derived foodstuffs and in animal feedstuffs should be favourably noted. These programmes should be coordinated nationwide.
Appendix V (contd)

Correlated results should be freely available on request to existing and prospective trading partner countries. Analytical methods and result reporting should be consistent with internationally recognised standards. If official responsibility for these programmes does not rest with the Veterinary Services, there should be appropriate provision to ensure that the results of such programmes are made available to the Veterinary Services for assessment.

4. Veterinary medicines

It should be acknowledged that primary control over veterinary medicinal products may not rest with the Veterinary Authorities in some countries, owing to differences between governments in the division of legislative responsibilities. However, for the purpose of evaluation, the Veterinary Services should be able to demonstrate the existence of effective controls (including nationwide consistency of application) over the manufacture, importation, export, registration, supply, sale and use of veterinary medicines, biologicals and diagnostic reagents, whatever their origin. The control of veterinary medicines has direct relevance to the areas of animal health and public health.

In the animal health sphere, this has particular application to biological products. Inadequate controls on the registration and use of biological products leave the Veterinary Services open to challenge over the quality of animal disease control programmes and over safeguards against animal disease introduction in imported veterinary biological products.

It is valid, for evaluation purposes, to seek assurances of effective government controls over veterinary medicines in so far as these relate to the public health risks associated with residues of these chemicals in animals and animal-derived foodstuffs. This process should be consistent with the standards set by the Codex Alimentarius or with alternative requirements set by the importing country where the latter are scientifically justified.

5. Integration between animal health controls and veterinary public health

The existence of any organised programme which incorporates a structured system of information feedback from inspection in establishments producing products of animal origin, in particular meat or dairy products, and applies this in animal health control should be favourably noted. Such programmes should be integrated within a national disease surveillance scheme.

Veterinary Services which direct a significant element of their animal health programmes specifically towards minimising microbial and chemical contamination of animal-derived products in the human food chain should receive favourable recognition in the evaluation. There should be evident linkage between these programmes and the official control of veterinary medicines and relevant agricultural chemicals.

Article 1.3.4.10.

Performance assessment and audit programmes

1. Strategic plans

The objectives and priorities of the Veterinary Services can be well evaluated if there is a published official strategic plan which is regularly updated. Understanding of functional activities is enhanced if an operational plan is maintained within the context of the strategic plan. The strategic and operational plans, if these exist, should be included in the evaluation.
Veterinary Services which use strategic and operational plans may be better able to demonstrate effective management than countries without such plans.

2. Performance assessment

If a strategic plan is used, it is desirable to have a process which allows the organisation to assess its own performance against its objectives. Performance indicators and the outcomes of any review to measure achievements against pre-determined performance indicators should be available for evaluation. The results should be considered in the evaluation process.

3. Compliance

Matters which can compromise compliance and adversely affect a favourable evaluation include instances of inaccurate or misleading official certification, evidence of fraud, corruption, or interference by higher political levels in international veterinary certification, and lack of resources and poor infrastructure.

It is desirable that the Veterinary Services contain (or have a formal linkage with) an independent internal unit/section/commission the function of which is to critically scrutinise their operations. The aim of this unit should be to ensure consistent and high integrity in the work of the individual officials in the Veterinary Services and of the corporate body itself. The existence of such a body can be important to the establishment of international confidence in the Veterinary Services.

An important feature when demonstrating the integrity of the Veterinary Services is their ability to take corrective action when miscertification, fraud or corruption has occurred.

A supplementary or an alternative process for setting performance standards and application of monitoring and audit is the implementation of formal quality systems to some or all activities for which the Veterinary Services are responsible. Formal accreditation to international quality system standards should be utilised if recognition in the evaluation process is to be sought.

4. Veterinary Services administration

a) Annual reports

Official government annual reports should be published, which provide information on the organisation and structure, budget, activities and contemporary performance of the Veterinary Services. Current and retrospective copies of such reports should be available to counterpart Services in other countries, especially trade partners.

b) Reports of government review bodies

The reports of any periodic or ad hoc government reviews of Veterinary Services or of particular functions or roles of the Veterinary Services should be considered in the evaluation process. Details of action taken as a consequence of the review should also be accessible.
Appendix V (contd)

c) Reports of special committees of enquiry or independent review bodies

    Recent reports on the *Veterinary Services* or elements of their role or function, and details of any subsequent implementation of recommendations contained in these reports should be available. The *Veterinary Services* concerned should recognise that the provision of such information need not be detrimental to the evaluation outcome; in fact, it may demonstrate evidence of an effective audit and response programme. The supplying of such information can reinforce a commitment to transparency.

d) In-service training and development programme for staff

    *In order to maintain a progressive approach to meeting the needs and challenges of the changing domestic and international role of Veterinary Services, the national administration should have in place an organised programme which provides appropriate training across a range of subjects for relevant staff.* This programme should include participation in scientific meetings of animal health organisations. Such a programme should be used in assessing the effectiveness of the Services.

e) Publications

    *Veterinary Services* can augment their reputation by demonstrating that their staff publish scientific articles in refereed veterinary journals or other publications.

f) Formal linkages with sources of independent scientific expertise

    Details of formal consultation or advisory mechanisms in place and operating between the *Veterinary Services* and local and international universities, scientific institutions or recognised veterinary organisations should be taken into consideration. These could serve to enhance the international recognition of the *Veterinary Services*.

g) Trade performance history

    In the evaluation of the *Veterinary Services* of a country, it is pertinent to examine the recent history of their performance and integrity in trade dealings with other countries. Sources of such historical data may include Customs Services.

| Article 1.3.4.11. |

**Participation in OIE activities**

Questions on a country's adherence to its obligations as a member of the OIE are relevant to an evaluation of the *Veterinary Services* of the country. Self-acknowledged inability or repeated failure of a Member Country to fulfil reporting obligations to the OIE will detract from the overall outcome of the evaluation. Such countries, as well as non-member countries, will need to provide extensive information regarding their *Veterinary Services* and sanitary/zoosanitary status for evaluation purposes.
Evaluation of the veterinary statutory body

1. **Scope**

   In the evaluation of the veterinary statutory body, the following items may be considered, depending on the purpose of the evaluation:

   - objectives and functions;
   - legislative basis, including autonomy and functional capacity;
   - human resources, including the composition and representation of the body’s membership;
   - institutional arrangements, accountability and transparency of decision-making;
   - sources and management of funding;
   - functional capabilities, including the ability to enforce its decisions (for example regarding registration requirements, standards of conduct, and disciplinary procedures);
   - administration of education training programmes and continuing professional development for veterinarians and veterinary para-professionals.

2. **Evaluation of objectives and functions**

   The veterinary statutory body should define its policy and objectives, including detailed descriptions of its powers and functions such as:

   - to regulate veterinarians and veterinary para-professionals through licensing and/or registration of such persons;
   - to determine the minimum standards of training required for degrees, diplomas and certificates entitling the holders thereof to be registered as veterinarians and veterinary para-professionals;
   - to determine the standards of professional conduct of veterinarians and veterinary para-professionals and to ensure these standards are met.

3. **Evaluation of legislative basis, autonomy and functional capacity**

   The veterinary statutory body should be able to demonstrate that it has the capacity, supported by appropriate legislation, to exercise and enforce control over all veterinarians and veterinary para-professionals. These controls should include, where appropriate, compulsory licensing and registration, minimum standards of training for the recognition of degrees, diplomas and certificates, setting standards of professional conduct and exercising control and the application of disciplinary procedures.
Appendix V (contd)

The veterinary statutory body should be able to demonstrate autonomy from undue political and commercial interests.

Where applicable, regional agreements for the recognition of degrees, diplomas and certificates for veterinarians and veterinary para-professionals should be demonstrated.

4. Evaluation of membership representation

Detailed descriptions should be available in respect of the membership of the veterinary statutory body and the method and duration of appointment of members. Such information includes:

- veterinarians designated by the Veterinary Administration, such as the Chief Veterinary Officer;
- veterinarians elected by members registered by the veterinary statutory body;
- veterinarians designated or nominated by the veterinary association(s);
- representative(s) of veterinary para-professions;
- representative(s) of veterinary academia;
- representative(s) of other stakeholders from the private sector;
- election procedures and duration of appointment;
- qualification requirements for members.

5. Evaluation of accountability and transparency of decision-making

Detailed information should be available on disciplinary procedures regarding the conducting of enquiries into professional misconduct, transparency of decision-making, publication of findings, sentences and mechanisms for appeal.

Additional information regarding the publication at regular intervals of activity reports, lists of registered or licensed persons including deletions and additions should also be taken into consideration.

6. Evaluation of financial sources and financial management

Information regarding income and expenditure, including fee structure(s) for the licensing/registration of persons should be available.

7. Evaluation of training programmes and programmes for continuing professional development, for veterinarians and veterinary para-professionals

Descriptive summary of continuing professional development, training and education programmes should be provided, including descriptions of content, duration and participants; documented details of quality manuals and standards relating to Good Veterinary Practice should be provided.
Appendix V (contd)

Article 1.3.4.13.

1. The Veterinary Services of a country may undertake self-evaluation against the above criteria for such purposes as national interest, improvement of internal efficiency or export trade facilitation. The way in which the results of self-evaluation are used or distributed is a matter for the country concerned.

2. A prospective importing country may undertake an evaluation of the Veterinary Services of an exporting country as part of a risk analysis process, which is necessary to determine the sanitary or zoosanitary measures which the country will use to protect human or animal life or health from disease or pest threats posed by imports. Periodic evaluation reviews are also valid following the commencement of trade.

3. In the case of evaluation for the purposes of international trade, the authorities of an importing country should use the principles elaborated above as the basis for the evaluation and should attempt to acquire information according to the model questionnaire outlined in Article 1.3.4.14. The Veterinary Services of the importing country are responsible for the analysis of details and for determining the outcome of the evaluation after taking into account all the relevant information. The relative ranking of importance ascribed, in the evaluation, to the criteria described in this Chapter will necessarily vary according to case-by-case circumstances. This ranking should be established in an objective and justifiable way. Analysis of the information obtained in the course of an evaluation study must be performed in as objective a manner as possible. The validity of the information should be established and reasonableness should be employed in its application. The assessing country must be willing to defend any position taken on the basis of this type of information, if challenged by the other party.

Article 1.3.4.14.

This Article outlines appropriate information requirements for the self-evaluation or evaluation of the Veterinary Services of a country.

1. Organisation and structure of Veterinary Services

   a) National Veterinary Services

      Organisational chart including numbers, positions and numbers of vacancies.

   b) Sub-national Veterinary Services

      Organisational charts including numbers, positions and number of vacancies.

   c) Other providers of Veterinary Services

      Description of any linkage with other providers of Veterinary Services.

2. National information on human resources

   a) Veterinarians

      i) Total numbers of veterinarians registered/licensed by the Veterinary statutory body of the country:
Appendix V (contd)

ii) Numbers of:
   - full time government veterinarians: national and sub-national;
   - part time government veterinarians: national and sub-national;
   - private veterinarians authorised by the Veterinary Services to perform official veterinary functions; [Describe accreditation standards, responsibilities and/or limitations applying to these private veterinarians.]
   - other veterinarians.

iii) Animal health:

Numbers associated with farm livestock sector on a majority time basis in a veterinary capacity, by geographical area [Show categories and numbers to differentiate staff involved in field service, laboratory, administration, import/export and other functions, as applicable.]:

   - full time government veterinarians: national and sub-national;
   - part time government veterinarians: national and sub-national;
   - other veterinarians.

iv) Veterinary public health:

Numbers employed in food inspection on a majority time basis, by commodity [Show categories and numbers to differentiate staff involved in inspection, laboratory and other functions, as applicable.]:

   - full time government veterinarians: national and sub-national;
   - part time government veterinarians: national and sub-national;
   - other veterinarians.

v) Numbers of veterinarians relative to certain national indices:

   - per total human population;
   - per farm livestock population, by geographical area;
   - per livestock farming unit, by geographical area.

vi) Veterinary education:

   - number of veterinary schools;
   - length of veterinary course (years);
   - international recognition of veterinary degree.

vii) Veterinary professional associations.
b) Graduate personnel (non-veterinary)

Details to be provided by category (including biologists, biometricians, economists, engineers, lawyers, other science graduates and others) on numbers within national Veterinary Services and available to national Veterinary Services.

c) Veterinary para-professionals employed by the Veterinary Services

i) Animal health:

- Categories and numbers involved with farm livestock on a majority time basis:
  - by geographical area;
  - proportional to numbers of field Veterinary Officers in the Veterinary Services, by geographical area.

- Education/training details.

ii) Veterinary public health:

- Categories and numbers involved in food inspection on a majority time basis:
  - meat inspection: export meat establishments with an export function and domestic meat establishments (no export function);
  - dairy inspection;
  - other foods.

- Numbers in import/export inspection.

- Education/training details.

d) Support personnel

Numbers directly available to Veterinary Services per sector (administration, communication, transport).

e) Descriptive summary of the functions of the various categories of staff mentioned above

f) Veterinary, veterinary para-professionals, livestock owner, farmer and other relevant associations

g) Additional information and/or comments.

3. Financial management information

a) Total budgetary allocations to the Veterinary Services for the current and past two fiscal years:

i) for the national Veterinary Services;

ii) for each of any sub-national veterinary authorities;

iii) for other relevant government-funded institutions.
Appendix V (contd)

b) Sources of the budgetary allocations and amount:

i) government budget;

ii) sub-national authorities;

iii) taxes and fines;

iv) grants;

v) private services.

c) Proportional allocations of the amounts in a) above for operational activities and for the programme components of Veterinary Services.

d) Total allocation proportionate of national public sector budget. [This data may be necessary for comparative assessment with other countries which should take into account the contexts of the importance of the livestock sector to the national economy and of the animal health status of the country.]

e) Actual and proportional contribution of animal production to gross domestic product.

4. Administration details

a) Accommodation

Summary of the numbers and distribution of official administrative centres of the Veterinary Services (national and sub-national) in the country.

b) Communications

Summary of the forms of communication systems available to the Veterinary Services on a nation-wide and local area bases.

c) Transport

i) Itemised numbers of types of functional transport available on a full-time basis for the Veterinary Services. In addition provide details of transport means available part-time.

ii) Details of annual funds available for maintenance and replacement of motor vehicles.

5. Laboratory services

a) Diagnostic laboratories (laboratories engaged primarily in diagnosis)

i) Descriptive summary of the organisational structure and role of the government veterinary laboratory service in particular its relevance to the field Veterinary Services.
ii) Numbers of veterinary diagnostic laboratories operating in the country:

- government operated laboratories;

- private laboratories accredited by government for the purposes of supporting official or officially-endorsed animal health control or public health testing and monitoring programmes and import/export testing.

iii) Descriptive summary of accreditation procedures and standards for private laboratories.

iv) Human and financial resources allocated to the government veterinary laboratories, including staff numbers, graduate and post-graduate qualifications and opportunities for further training.

v) List of diagnostic methodologies available against major diseases of farm livestock (including poultry).

vi) Details of collaboration with external laboratories including international reference laboratories and details on numbers of samples submitted.

vii) Details of quality control and assessment (or validation) programmes operating within the veterinary laboratory service.

viii) Recent published reports of the official veterinary laboratory service which should include details of specimens received and foreign animal disease investigations made.

ix) Details of procedures for storage and retrieval of information on specimen submission and results.

x) Reports of independent reviews of the laboratory service conducted by government or private organisations (if available).

xi) Strategic and operational plans for the official veterinary laboratory service (if available).

b) Research laboratories (laboratories engaged primarily in research)

i) Numbers of veterinary research laboratories operating in the country:

- government operated laboratories;

- private laboratories involved in full time research directly related to animal health and veterinary public health matters involving production animal species.

ii) Summary of human and financial resources allocated by government to veterinary research.

iii) Published programmes of future government sponsored veterinary research.

iv) Annual reports of the government research laboratories.
Appendix V (contd)

6. Functional capabilities and legislative support

a) Animal health and veterinary public health

i) Assessment of the adequacy and implementation of relevant legislation (national or sub-national) concerning the following:

- animal and veterinary public health controls at national frontiers;
- control of endemic animal diseases, including zoonoses;
- emergency powers for control of exotic disease outbreaks, including zoonoses;
- inspection and registration of facilities;
- veterinary public health controls of the production, processing, storage and marketing of meat for domestic consumption;
- veterinary public health controls of the production, processing, storage and marketing of fish, dairy products and other foods of animal origin for domestic consumption;
- registration and use of veterinary pharmaceutical products including vaccines.

ii) Assessment of ability of Veterinary Services to enforce legislation.

b) Export/import inspection

i) Assessment of the adequacy and implementation of relevant national legislation concerning:

- veterinary public health controls of the production, processing, storage and transportation of meat for export;
- veterinary public health controls of production, processing, storage and marketing of fish, dairy products and other foods of animal origin for export;
- animal health and veterinary public health controls of the export and import of animals, animal genetic material, animal products, animal feedstuffs and other products subject to veterinary inspection;
- animal health controls of the importation, use and bio-containment of organisms which are aetiological agents of animal diseases, and of pathological material;
- animal health controls of importation of veterinary biological products including vaccines;
- administrative powers available to Veterinary Services for inspection and registration of facilities for veterinary control purposes (if not included under other legislation mentioned above);
- documentation and compliance.

ii) Assessment of ability of Veterinary Services to enforce legislation.
Appendix V (contd)

7. **Animal health and veterinary public health controls**

a) **Animal health**

   i) Description of and sample reference data from any national animal disease reporting system controlled and operated or coordinated by the *Veterinary Services*.

   ii) Description of and sample reference data from other national animal disease reporting systems controlled and operated by other organisations which make data and results available to *Veterinary Services*.

   iii) Description and relevant data of current official control programmes including:

      - epidemiological surveillance or monitoring programmes;
      - officially approved industry administered control or eradication programmes for specific diseases.

   iv) Description and relevant details of animal disease emergency preparedness and response plans.

   v) Recent history of animal disease status:

      - animal diseases eradicated nationally or from defined sub-national *zones* in the last ten years;
      - animal diseases of which the prevalence has been controlled to a low level in the last ten years;
      - animal diseases introduced to the country or to previously free sub national regions in the last ten years;
      - emerging diseases in the last ten years;
      - animal diseases of which the prevalence has increased in the last ten years.

b) **Veterinary public health**

   i) **Food hygiene**

      - Annual national slaughter statistics for the past three years according to official data by species of animals (bovine, ovine, porcine, caprine, poultry, farmed game, wild game, equine, other).

      - Estimate of total annual slaughterings which occur but are not recorded under official statistics.

      - Proportion of total national slaughter which occurs in registered export establishments, by category of animal.
Appendix V (contd)

- Proportion of total national slaughter which occurs under veterinary control, by category of animal.

- Numbers of commercial fresh meat establishments in the country which are registered for export by national Veterinary Services:
  - slaughterhouses (indicate species of animals);
  - cutting/packing plants (indicate meat type);
  - meat processing establishments (indicate meat type);
  - cold stores.

- Numbers of commercial fresh meat establishments in the country approved by other importing countries which operate international assessment inspection programmes associated with approval procedures.

- Numbers of commercial fresh meat establishments under direct public health control of the Veterinary Services (including details of category and numbers of inspection staff associated with these premises).

- Description of the veterinary public health programme related to production and processing of animal products for human consumption (including fresh meat, poultry meat, meat products, game meat, dairy products, fish, fishery products, molluses and crustaceans and other foods of animal origin) especially including details applying to exports of these commodities.

- Descriptive summary of the roles and relationships of other official organisations in public health programmes for the products listed above if the national Veterinary Services do not have responsibility for those programmes which apply to national production destined to domestic consumption and/or exports of the commodities concerned.

ii) Zoonoses

- Descriptive summary of the numbers and functions of staff of the Veterinary Services involved primarily with monitoring and control of zoonotic diseases.

- Descriptive summary of the role and relationships of other official organisations involved in monitoring and control of zoonoses to be provided if the national Veterinary Services do not have these responsibilities.

iii) Chemical residue testing programmes

- Descriptive summary of national surveillance and monitoring programmes for environmental and chemical residues and contaminants applied to animal-derived foodstuffs, animals and animal feedstuffs.
Appendix V (contd)

- Role and function in these programmes of the national *Veterinary Services* and other *Veterinary Services* to be described in summary form.

- Descriptive summary of the analytical methodologies used and their consistency with internationally recognised standards.

iv) Veterinary medicines

- Descriptive summary of the administrative and technical controls involving registration, supply and use of veterinary pharmaceutical products especially including biological products. This summary should include a focus on veterinary public health considerations relating to the use of these products in food-producing animals.

- Role and function in these programmes of the national *Veterinary Services* and other *Veterinary Services* to be described in summary form.

8. Quality systems

   a) Accreditation

   Details and evidence of any current, formal accreditation by external agencies of the *Veterinary Services* of any components thereof.

   b) Quality manuals

   Documented details of the quality manuals and standards which describe the accredited quality systems of the *Veterinary Services*.

   c) Audit

   Details of independent (and internal) audit reports which have been undertaken of the *Veterinary Services* of components thereof.

9. Performance assessment and audit programmes

   a) Strategic plans and review

   i) Descriptive summary and copies of strategic and operational plans of the *Veterinary Services* organisation.

   ii) Descriptive summary of corporate performance assessment programmes which relate to the strategic and operational plans - copies of recent review reports.

b) Compliance

   Descriptive summary of any compliance unit which monitors the work of the *Veterinary Services* (or elements thereof).
Appendix V (contd)

c) Annual reports of the national Veterinary Services

Copies of official annual reports of the national (sub-national) *Veterinary Services*.

d) Other reports

i) Copies of reports of official reviews into the function or role of the *Veterinary Services* which have been conducted within the past three years.

ii) Descriptive summary (and copy of reports if available) of subsequent action taken on recommendations made in these reviews.

e) Training

i) Descriptive summary of in-service and development programmes provided by the *Veterinary Services* (or their parent Ministries) for relevant staff.

ii) Summary descriptions of training courses and duration.

iii) Details of staff numbers (and their function) who participated in these training courses in the last three years.

f) Publications

Bibliographical list of scientific publications by staff members of *Veterinary Services* in the past three years.

g) Sources of independent scientific expertise

List of local and international universities, scientific institutions and recognised veterinary organisations with which the *Veterinary Services* have consultation or advisory mechanisms in place.

10. Membership of the OIE

State if country is a member of the OIE and period of membership.

11. Other assessment criteria

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* OIE Terrestrial Animal Health Standards Commission/September 2005
Appendix VI

Performance, Vision and Strategy (PVS) for

VETERINARY SERVICES (VS)¹

Introduction

In this era of globalization, the development and growth in many countries depends on the performance of their agricultural economies, and this, in turn, directly relates to the quality of their national veterinary services (VS). VS play also a major role in Veterinary public health including food-borne diseases and regional and international market access for animals and their products. To be effective, VS should operate based on scientific principles and be technically independent and immune from political pressures of its users’. However, efforts to strengthen official services, requires the active participation and investment on the part of both the public and the private sectors. To assist in this effort, the World Organization for Animal Health (OIE) and the Inter-American Institute for Cooperation on Agriculture (IICA) have joined forces to develop the Performance, Vision and Strategy (PVS) instrument. The PVS instrument can assist VS to establish their current level of performance, form a shared vision with the private sector, establish priorities and facilitate strategic planning in order to take full advantage of the new opportunities and obligations of globalization.

The OIE promotes animal health and public health including food-borne diseases safety in the international trade of animals and their related products by issuing harmonized sanitary guidelines on international certification and disease control methods and working to improve the resources and legal framework of the VS. Likewise, IICA helps to strengthen VS so they can be more efficient and competitive nationally and internationally and can contribute to the improved health of their consumers. Both organizations share a mutual interest to help countries comply with the Agreement on the Application of Sanitary and Phytosanitary Measures (SPS) of the World Trade Organization (WTO) and the standards, guidelines and recommendations of the OIE.

The traditional mission of VS has been to protect domestic agriculture and, over time, most of its resources were channeled toward the control of diseases² that threatened primary production. The focus of the services provided were from the national borders inward and the credibility of these services, in the eyes of its users and other countries, depended in large measure on the effectiveness of its domestic programs, and its response to emergencies arising from the entry of foreign diseases.

In light of the growing international requirements and opportunities facing countries, it behooves VS to adopt a broader mandate and vision, and provide new services that complement the portfolio of existing services. This will entail stronger alliances and closer cooperation with its users, other countries and their national veterinary service counter parts. The WTO/SPS agreement reaffirms the right of the member countries to protect plant, animal and human life or health, but the agreement also requires that countries base their SPS measures on scientific principles and the OIE standards - the fundamental basis of operation to ensure that international trade is free of discrimination and scientifically unjustified restrictions.

¹ Veterinary services means the Veterinary Administration, all the Veterinary Authorities, and all persons authorized, registered or licensed by the veterinary statutory body of a country. They will be called “VS” in all the document

² Clinical and/or pathological manifestation of an infection
Appendix VI (contd)

Experience has shown that those countries, whose VS are more developed and credible in the eyes of its users, trading partners and other countries, contain four fundamental components: 1) the technical capability to address current and new issues based on scientific principles; 2) the human and financial capital to attract resources and retain professionals with technical and leadership skills; 3) the interaction with the private sector in order to stay on course and carry out relevant joint programs and services; and 4) the ability to access markets through the compliance with existing standards and the implementation of new disciplines such as harmonization of standards, equivalence and regionalization. These four components provide the basic structure of the PVS instrument.

Applying the PVS Instrument

To establish the current level of performance, form a shared vision, establish priorities and facilitate strategic planning, a series of five to eight critical competencies have been developed for each of the four fundamental components. For each critical competency, qualitative levels of advancement are described. To help visualize the potential or cumulative level of advancement within each critical competency, a pie chart is shown next to the written explanation for each level. A higher level of advancement assumes that the VS is complying with the preceding (and non zero) levels.

In addition to the qualitative levels, additional space has been provided after each critical competency to expand upon or clarify responses, if so desired. The following hypothetical example illustrates the level of advancement determined along with an explanation for the critical competency harmonization, one of the [twenty-eight] critical competencies in the PVS instrument.

3. Harmonization

The capability and authority of the VS to be active in harmonization and ensure that the national regulatory norms covered under its mandate are in conformity with relevant international standards, guidelines and recommendations.

Levels of advancement:

0. The VS has no process to be aware of international standards. National regulatory norms do not take account of international standards, guidelines and recommendations.
1. The VS is aware of relevant standards but has no process to identify gaps, inconsistencies, or non-conformities in national regulatory norms as compared to international standards, guidelines and recommendations.
2. The VS monitors the establishment of new international standards, guidelines and recommendations and periodically reviews national regulatory norms with the aim of harmonizing them as appropriate with international standards, guidelines and recommendations.
3. Same as previous level plus the VS is active in reviewing and commenting on draft standards, guidelines and recommendations.
4. Same as previous level plus the VS actively and regularly participates at the international level in the formulation of international standards, guidelines and recommendations.*

* A country could be active in international standard setting without actively pursuing national changes. The importance of this element is to promote national change.
Using the results

The PVS instrument is designated for easy understanding and is flexible in its application and use. More than a diagnostic tool, it is a process oriented towards the future which can be used in passive or active mode, depending on the level of interest and commitment by the users and the official service in improving their national services over time.

If it is used in the passive mode, the PVS instrument raises awareness, improves understanding and guides the different sectors participating in the process regarding the basic components and critical competencies the VS must contain in order to function adequately. In this mode the instrument can also be used to develop a shared vision, foster dialogue and adopt a common language for discussion.

The active mode is where the maximum potential is generated and the best results can be obtained, assuming the commitment is present on the part of both the public and private sector. In this mode, performance is assessed, differences are explored and priorities are established. Leadership on the part of the public sector is a critical element for success. This active mode is where actions happen, investments are evaluated and made and commitment is carried out. Continuity of the PVS process is assured when a true partnership between the official and the private sector exists.

As a very important additional reference, Chapter 1.3.3 on the Evaluation of Veterinary Services, in the Terrestrial Animal Health Code of the OIE and Chapter 1.4.3 on the Evaluation of Competent Authorities, in the Aquatic Animal Health Code, expand upon and further clarifies some of the levels of advancement described in some of the critical competencies of the PVS instrument. The instrument can be used to facilitate the dialogue with different users in the public and private sectors that share a common interest in improving the vision and performance of the public services. For example, the interested parties can jointly participate in establishing the current level of performance, identifying priorities and adopting actions that strengthen the national services. In addition, the director of the national VS can use the instrument to monitor progress in each one of the four components.

For the VS, the results of the PVS instrument can help to: 1) indicate the overall performance of each one of the four components; 2) rate the relative performance within each one of the critical competencies; 3) compare the performance of the VS with that of other veterinary services in the region or globally, in order to explore areas for cooperation or negotiation; 4) identify the differences in the responses of the different users in order to arrive at common points of view; 5) foster common understanding in order to achieve greater levels of advancement; 6) help determine the benefits and costs of investing in VS and obtaining assistance from financial and technical cooperation agencies, 7) provide a basis for establishing a routine monitoring and follow up mechanism on the overall level of performance of the VS over time; and 8) help identify and present objectives and specific needs when applying for financial support (loans and/or grants). 9) Prepare a process of verification of compliance with OIE standards on quality and evaluation of VS by an external independent body under the auspices of the OIE.

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3 OIE standards allow importing countries to make audits in exporting countries and in particular check the compliance of exporting countries with OIE standards on quality and evaluation of VS
FUNDAMENTAL COMPONENTS

I. TECHNICAL CAPABILITY
II. HUMAN AND FINANCIAL CAPITAL
III. INTERACTION WITH THE PRIVATE SECTOR
IV. ACCESS TO MARKETS
I. TECHNICAL CAPABILITY

The capability of the VS to establish and apply sanitary measures and science-based procedures.

Critical competencies:

1. Diagnostic capability
2. Early detection and emergency response capability
3. Quarantine
4. Epidemiological surveillance
5. Quality systems
6. Risk analysis
7. Technical innovation

1. Diagnostic capability

The capability and authority of the VS to identify and record those biological, physical and chemical agents including those relevant for public health that can adversely affect animals and their related products.

Levels of advancement:

0. For existing diseases, the VS can carry out the clinical diagnosis, but not the laboratory\textsuperscript{4} confirmation.

1. For zoonoses\textsuperscript{5} and other diseases with a major economic or public health impact, the VS can collect samples in the country and immediately ship them to the laboratory for confirmation.

2. For zoonoses, and other diseases not present in the country, but known to exist in the region or could enter via trade, the VS has procedures in place to collect samples and immediately ship them to the laboratory for confirmation.

3. In the case of new and emerging diseases in the region or world, the VS has access to a network of national or international reference laboratories and can collect and ship samples to the most qualified laboratory for confirmation.

4. The VS actively promotes the accreditation of its laboratories and audits\textsuperscript{6} the quality of its clinical diagnostic, collection and shipment of samples procedures.

2. Early detection and emergency response capability

The capability and authority of the VS to rapidly respond to unexpected disease outbreak\textsuperscript{7} or other situations that put at immediate risk the sanitary status\textsuperscript{8} of the animal populations covered under its mandate.

\textsuperscript{4} Means a properly equipped institution staffed by technically competent personnel under the control of a specialist in veterinary diagnostic methods, who is responsible for the validity of the results. The Veterinary Administration approves and monitors such laboratories with regard to the diagnostic tests required for international trade.

\textsuperscript{5} Zoonoses (Zoonotic diseases): Any disease or infection which is naturally transmissible from animals to humans.

\textsuperscript{6} Audits: A systematic and functionally independent examination, the objective of which is to determine if an activity or process and subsequent results meet the prescribed objectives.

\textsuperscript{7} Outbreak means an occurrence of one of the diseases listed by the OIE in an establishment, breeding establishment or premises, including all buildings and all adjoining premises, where animals are present. Where it cannot be defined in this way, the outbreak shall be considered as occurring in the part of the territory in which, taking local conditions into account, it cannot be guaranteed that both susceptible and non-susceptible animals have had no direct contact with affected or suspected cases in that area.
Appendix VI (contd)

Levels of advancement:

0. The VS has no field network nor system to determine whether or not a sanitary emergency exists and it does not have the authority to declare such an emergency and take action.

1. The VS has a field network and a system to determine whether or not a sanitary emergency exists but lacks the necessary legal and financial support\(^9\) to take action in response to sanitary emergencies.

2. The VS has a system to make timely decisions on whether or not a sanitary emergency exists. The VS has the legal framework and funding sources to take action in response\(^10\) to sanitary emergencies through an efficient national chain of command.

3. Same as previous level plus the VS has contingency plans or general action plans for diseases of concern that enable it to coordinate actions with other relevant organizations or institutions and the private sector (including veterinary practitioner), in response to sanitary emergencies through an efficient national chain of command.

3. **Quarantine**

The capability and authority of the VS to prevent the entrance and spread of unwanted diseases in the country.

Levels of advancement:

0. The VS does not compile information on the sanitary status in its own country or maintain any type of quarantine procedures with its neighbouring countries or trading partners.

1. The VS has up-to-date information on exporting countries which it incorporates into its quarantine procedures for the commercial trade of primarily farm animals and their related products that come into the country and may threaten its sanitary status.

2. The VS has up-to-date information on exporting countries which it incorporates into quarantine procedures for animals and their related products, even if of no significant trade or commercial value (e.g. companion animals) but enter into the country through established trade channels.

3. The VS can or has implemented specialized quarantine programs\(^11\) in the country of origin for specific animals and their related products.

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\(^8\) The status of a country or compartment within the country with respect to a particular disease, in accordance to the criteria set forward in the Terrestrial Animal Health Code of the OIE.

\(^9\) The phrase, legal and financial support, refers to the VS already having in place the legal framework and financial resources in order to take immediate actions.

\(^10\) Appropriate response to sanitary emergency includes an appropriate early detection system.

\(^11\) Programs that facilitate the detection of transmissible diseases and make it possible to evaluate the health of the population in question before being transported.
4. The VS carries out quality assurance audits of its own quarantine procedures and, if necessary, those of its trading partners, in compliance with OIE standards on quality and evaluation of VS.

4. **Epidemiological surveillance**

The capability and authority of the VS to determine, monitor and verify the sanitary status of the animal populations covered under its mandate.

**Levels of advancement:**

0. The VS has no program in place for surveillance or monitoring.

1. The VS conducts a surveillance program based on existing information or suspected cases, where samples are collected and sent to the laboratories.

2. The VS conducts active monitoring programs in animal populations on diseases of economic and zoonotic importance.

3. The VS conducts surveillance programs in populations of greatest risk covering zoonoses, and other diseases of economic importance.

4. The VS structures its surveillance programs taking into account the sanitary status of its neighboring countries and trade flows.

5. **Quality systems**

The authority and capacity of VS to define their veterinary public health policies, formalize their activities, in particular concerning control and certification and making sure that these are well executed.

**Levels of advancement:**

0. The VS has no system for the control of their activities.

1. The VS has established an administrative structure capable of ensuring the chain of command, defining the required regulations and delegation of authority.

2. The VS has defined the policies and has evaluated the resource needs.

3. The VS has implemented a general system for registering their procedures and instructions.

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12 The term, surveillance, refers to the ongoing and systematic process of collecting, analyzing, interpreting and disseminating information on the sanitary status, including early detection of exotic and emerging diseases. The term, monitoring, is more specific in its application and is directed at detecting changes in the prevalence of a pest or disease for a given population and environment. Surveillance and monitoring procedures take into account as a minimum basis the requirements published in the appendices of the relevant chapters of the OIE *Codes and Manuals*. 

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Appendix V (contd)

4. The VS has a system for the evaluation of the effectiveness of their services (internal audit).

5. The VS is subjected to external audits of its Quality system.

6. Risk analysis\textsuperscript{13}

The capability of the VS to make decisions and carry out actions based on scientific principles and evidence, including the assessment, communication and management of risk.

Levels of advancement:

0. The VS does not compile data or other kinds of information that could be used to identify potential sanitary hazards and analyze risks. Sanitary decisions are not supported by scientific evidence.

1. The VS compiles and maintains sources of information or can access the information necessary in order to assess risks. Sanitary decisions may be based on scientific evidence.

2. The VS has a system to actively seek and maintain relevant data and information for risk assessment and dedicated personnel with this responsibility. Scientific principles and evidence provide the basis for options considered by sanitary decision makers in order to manage risks.

3. Same as previous level plus the VS is consistent in conducting scientifically based risk assessments in compliance with relevant OIE standards and communicating the decisions taken to the WTO/SPS, the OIE and its relevant trading partners.

4. Same as previous level plus the VS is consistent in managing and communicating the risks in conformity with the WTO/SPS Agreement and relevant standards of the OIE.

7. Technical innovation

The capability of the VS to update its overall service, in accordance with the latest scientific advances and based on the sanitary norms and measures of the OIE, Codex Alimentarius and the WTO/SPS Agreement.

Levels of advancement:

0. The VS has only informal access to technical innovations through personal contacts or external media sources.\textsuperscript{14}

1. The VS maintains information base on technical innovations and international norms through subscriptions to scientific journals and electronic media.

2. The VS carries out a specific program that identifies technical innovations which can improve its operation and procedures.

\textsuperscript{13} The term, risk, refers to the likelihood of an adverse event and the probable magnitude of the consequences in the importing country during a specified time period. Risk analysis refers to the assessment, management and communication of risk, not only for imports but for domestic issues which may also arise.

\textsuperscript{14} External media are those sources of information that may not be available or subscribed to by the VS such as scientific publications and magazines.
3. The VS incorporates technical innovations into selected functions and procedures, with specific resources and the collaboration or contributions of its users.\(^\text{15}\)

4. The VS has a dedicated budget plus the collaboration and contributions of its users, to continually implement technical innovations throughout the national service.

**II. HUMAN AND FINANCIAL CAPITAL**

Institutional and financial sustainability as evidenced by the level of professional talent and financial resources available.

Critical competencies:

1. Human talent
2. Training
3. Funding sources
4. Stability of policies and programs
5. Contingency funds
6. Technical independence
7. Capability to invest and grow

1. **Human talent (Initial training)**

The capability of the VS to efficiently carry out the professional and technical functions; measured in two ways: academic degrees\(^\text{16}\) and qualifications of its professional staff.

A veterinary positions:

Levels of advancement:

0 In the core of the VS the majority of the veterinary positions are not occupied by personnel holding a university diploma.

1 In the core of the VS the veterinary positions are defined in terms of the area of expertise, the placement within the structure, and the level of competence and of initial training (university degree recognized by the State).

2 In the core of the VS there is a service in charge of the management of human resources and of the appropriateness of positions and diplomas according to international standards.

3 The management of veterinary human resources is subject to internal audits.

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\(^\text{15}\) This includes consulting with the OIE, WTO, Codex websites and books for publications and notices and regular participation in international forum

\(^\text{16}\) Not all professional positions require a academic degree. Nonetheless, the rate of academic degrees serves as an indicator of the professional excellence within the VS.
Appendix VI (contd)

A technical and administrative positions:

Levels of advancement:

0. In the core of the VS the majority of technical and administrative positions are not occupied by personnel with professional qualifications\(^{17}\).

1. In the core of the VS the majority of technical and administrative positions are occupied by personnel with professional qualifications.

2. In the core of the VS the technical and administrative positions are defined in terms of the area of expertise, the placement within the structure, and the level of competence and of initial training (university degree recognized by the State).

3. In the core of the VS there is a service in charge of the management of human resources and of the appropriateness of positions and diplomas according to international standards.

4. The management of the entire human resources is subject to internal audits.

2. Training (Continuing education)

The capability of the VS to keep its personnel up-to-date in terms of relevant information and knowledge; measured in terms of the implementation of an annual training plan

Levels of advancement:

0. The VS has no training plans. (Continuing education plan)

1. The VS has training plans but they are not updated or funded.

2. The VS has annual training plans that are updated and funded but only partially implemented\(^{18}\).

3. The VS has updated and funded training plans largely implemented.

4. The VS has up to date training plans implemented for everyone.

3. Funding sources

The ability of the VS to access financial resources for its continued operation and sustainability, independent of any type of political pressure from users.

\(^{17}\) OIE international standards on quality and evaluation of VS make reference to the quality of the professional judgment.

\(^{18}\) Partially implemented may be only implemented for some personnel or only partially implemented for all personnel.
Appendix VI (contd)

Levels of advancement:

0. Funding for the VS is neither stable nor clearly defined. The budget for the national veterinary service competes with other State institutions and depends on resources allocated irregularly from the general treasury and/or non national donors.

1. The VS is funded from a continuous specific line item prescribed within the national budget as well as resources coming from non national donors if it is the case.

2. The VS is funded from a continuous specific line item prescribed within the national budget and with user fees generated by providing specific services (e.g. quarantine and certification services).

3. In addition to the previous levels, the VS also receives additional resources from its users to execute specific programs under complete transparency and ensuring full independence.

4. Stability of policies and programs

The capability of the VS to implement and sustain policies and programs over time; measured by the frequency of which the entire VS is reorganized and by the coordination capability between government institutions.

A. Levels of advancement (VS reorganization):

0. The VS is reorganized frequently at all levels.

1. The VS is reorganized frequently at some levels.

2. The VS is reorganized only at political levels after political changes.

3. The VS is reorganized only occasionally at political levels after political changes.

4. The VS is stable at technical and political levels.

B. Levels of advancement (coordination capability between government institutions):

0. The national regulations do not clearly define the obligations and competencies of all the official sector institutions that comprise the VS.

1. There are national regulations that define the obligations and competencies of the official sector institutions at the national and local levels.

2. There are coordinated inter and intra institutional activities in the official sector at least at the national level.

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19 Users means farmers, livestock traders and/or industry

20 In compliance with OIE international standards on quality regarding independency and impartiality.

21 A stable organization maintains its core structure and functions for 5 years or more

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Appendix VI (contd)

3. There are coordinated inter and intra institutional activities in the official sector at both the national and local levels.

5. Contingency funds

The capability of the VS to access extraordinary financial resources in order to respond to emergency situations or emerging issues; measured by the ease of which contingency resources can be made available.

Levels of advancement:

0. No contingency fund exists and any extraordinary resources can only be obtained through legislation or presidential decree.
1. A contingency fund with limited resources has been established, but any additional resources must be approved via presidential decree or law.
2. A contingency fund with limited resources has been established, but any additional resources must be approved by the Minister of Agriculture.
3. A contingency fund with substantial resources has been established, but additional resources must be approved by the Minister of Agriculture.
4. A contingency fund with substantial resources has been established and includes additional resources previously made available by its users\(^\text{22}\).

6. Technical independence

The capability of the VS to carry out its duties with autonomy and free from political interference that may affect technical and scientific decisions; measured in two ways: political appointments\(^\text{21}\) and technical support for decisions.

A. Levels of advancement (management positions):

0. The Director General of the entire agricultural health and food safety institution (if applicable), the Director of the VS and his/her direct reports are political appointees.
1. The Director General of the entire agricultural health and food safety institution (if applicable) and the Director of the VS are the only political appointees.
2. The selection of the Directors is not made only on political considerations.

B. Levels of advancement (technical support for decisions):

0. The technical decisions made by the VS are almost always based on political considerations.
1. The technical decisions incorporate scientific principles, but must be modified to conform to any political considerations.

\(^{22}\) “Users” means there all beneficiaries of the activities of VS, such as farmers, traders, consumers and industry.

\(^{21}\) The phrase, political appointments, refers to appointments made by the party in office, serving at the pleasure of politicians and subject to immediate removal.
2. The technical decisions are based on scientific principles but are subject to review and possible modification based on political considerations.

3. The technical decisions are based only on scientific principles and are not changed to meet any political considerations.\(^\text{24}\).

7. **Capability to invest and grow**

   The capability of the VS to secure additional investments over time that leads to a sustained improvement in the entire service. The utilization of such resources is not subject to any type of political pressure from its users.

   **Levels of advancement:**

   0. There are no sustained actions to support the overall structure of the VS.

   1. The VS elaborates and presents proposals and secures investment resources for improvements and infrastructures from cooperation or donor agencies.

   2. The VS secures over time, significant investment resources for improvements and infrastructure, through extraordinary allocations from the national (general treasury) or local public resources or special line items.

   3. In addition to the previous levels, the beneficiaries including farmers and/or industry provide resources to the VS for improvements and infrastructure\(^\text{25}\).

**III. INTERACTION WITH THE BENEFICIARIES**

The capability of the VS to collaborate with and involve the beneficiaries (including farmers and/or industry) in the implementation of programs and activities.

**Critical competencies:**

1. Communication
2. Consultation of beneficiaries
3. Official representation
4. Accreditation
5. Statutory body
6. Joint action programs implementation

1. **Communication**

   The capability of the VS to inform, in a transparent, effective and timely fashion, its users of activities, programs and developments.

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\(^{24}\) In accordance with the principles of the OIE *Codes* on quality of VS

\(^{25}\) In compliance with OIE standards on independence and impartiality of VS
Appendix VI (contd)

Levels of advancement:

0. The VS has no mechanism in place to keep users informed of activities, programs and sanitary developments.

1. The VS maintains an official communication outlet, which users can consult regarding standards, regulations and notifications.

2. The VS routinely\(^\text{26}\) publishes the results of its activities, programs and sanitary developments.

3. The VS provides up-to-date information, accessible via the internet, on sanitary developments and its programs and activities currently underway, and actively seeks input from the private sector, including farmers.

2. Consultation of beneficiaries

The capability of the VS to maintain fluid channels of consultation with the public and private sectors\(^\text{27}\) and users\(^\text{28}\).

Levels of advancement:

0. The VS has no consultation mechanisms in place to facilitate the dialogue between the relevant State institutions and the users.

1. The VS maintains informal channels of consultation with the relevant State institutions and the users.

2. The VS establishes and promotes official dialogue with the different users on its proposed and current regulations.

3. The VS holds forums and meetings with the different users in order to establish or improve its programs and services.

4. The VS actively promotes dialogue with and solicits feedback from the different users regarding national laws and regulations and official representation at the WTO/SPS and OIE.

5. The VS actively promotes dialogue with and solicits feedback from the different users regarding national laws and regulations and official representation at the WTO/SPS, OIE and Codex Alimentarius.

3. Official representation

The capability of the VS to regularly and actively participate, coordinate and provide follow up to the meetings of international organizations such as the WTO/SPS, OIE and Codex Alimentarius\(^\text{29}\).

\(^{26}\) Means every six months

\(^{27}\) private sector includes farmers, industry, transport and distribution

\(^{28}\) “users” means all beneficiaries of the VS activities

\(^{29}\) in compliance with international procedures and practices.
Levels of advancement:

0. The VS does not participate in or follow up on the meetings of the WTO/SPS, OIE and Codex Alimentarius.

1. The VS participates sporadically or passively\(^\text{30}\) in the meetings of the WTO/SPS, OIE and Codex Alimentarius.

2. The VS takes into consideration the opinions of its users and participates regularly and actively\(^\text{31}\) in the meetings of the WTO/SPS, OIE and Codex Alimentarius.

3. The VS, in consultation with its different users, identifies strategic topics, provides leadership and coordinates between the national delegations these topics over time as part of the agenda in the meetings of the WTO/SPS, OIE and Codex Alimentarius.

4. **Accreditation / Delegation**

   The capability and authority of the VS to accredit and delegate\(^\text{32}\) with third parties (e.g. private veterinarians, laboratories, etc), the execution of specific official services.

   Levels of advancement:

   0. The VS has neither the authority nor the capability to accredit and delegate to third parties.

   1. The VS has authority to accredit and delegate to third parties but no specific accreditation or delegation activities.

   2. The VS has accreditation and delegation programs for third parties and selected services.

   3. The VS can develop and implement accreditation and delegation programs for new services.

   4. The VS carries out quality assurance audits of its accreditation and delegation programs through an efficient national chain of command in order to maintain the trust of its trading partners.

5. **Statutory body**

   The veterinary statutory body, in accordance with the OIE’s definition, is an independent authority charged with the registration/licensing of veterinarians and authorization of veterinary para-professionals. Among others, it verifies the validity and the level of the veterinary diploma required to exercise the veterinary profession.

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\(^{30}\) *Passive participation* refers to being present at, but contributing little, to the meetings in question

\(^{31}\) *Active participation* refers to preparation in advance of, and contributing during the meetings in question, including exploring common solutions and generating proposals and compromises for possible adoption.

\(^{32}\) In compliance with OIE standards on quality of VS
Appendix VI (contd)

Levels of advancement:

0. There is no veterinary statutory body in the country.

1. There is a veterinary statutory body, but it does not have the power to discipline or make decisions.

2. The veterinary statutory body can only exercise its authority within the private sector.

3. The veterinary statutory body can also exercise its authority within the public sector.

4. The veterinary statutory body is subjected to auditing and evaluation procedures.

6. Joint programmes implementation

The capability of the VS and the private sector to formulate and implement joint programs on annual and/or pluri-annual bases.

Levels of advancement:

0. The VS has no joint programs.

1. The VS has established annual and/or pluri-annual joint programs but they are not updated or funded.

2. The VS has annual and/or pluri-annual joint programs that are updated and funded but only partially implemented.

3. The veterinary has joint programs that are updated annually and fully implemented.

IV. ACCESS TO MARKETS

The capability and authority of the VS to provide support in order to access, expand and retain regional and international markets for animals and animal products.

Critical competencies:

1. Compliance with regulations
2. Setting of regulations
3. Harmonization
4. Certification
5. Equivalency agreements
6. Traceability
7. Transparency
8. Zoning
9. Compartmentalization

33 Partially implemented may be only implemented for some activities or only partially implemented for all activities.
Appendix VI (contd)

1. **Compliance with regulations**

The capability and authority of the VS to ensure that users are in compliance with laws and regulations covered under its mandate.

Levels of advancement:

0. The VS has no program to ensure user compliance with laws and regulations.

1. The VS implements a compliance program consisting of inspection and verification of laws and regulations respect for selected animals, animal-products and processes, but only reports instances of non-compliance.

2. The VS implements a compliance program consisting of inspection and verification of laws and regulations respect for selected animals and animal products and processes, and, if necessary, imposes appropriate penalties in instances of non-compliance.

3. The VS implements a compliance program consisting of inspection and verification of laws and regulations respect for all animals, animal-products and processes covered under its mandate, and, if necessary, impose appropriate penalties in instances of non-compliance.

4. The VS carries out audits of its inspection and verification compliance programs through an efficient national chain of command.

2. **Setting of regulations**

The capability and authority of the VS to propose laws and to formulate and adopt regulations for animals, animal-products and processes covered under its mandate.

Levels of advancement:

0. The VS does not have the authority to prepare national legislation and set regulations.

1. The VS has the technical capability to propose national legislation and formulate regulations.

2. The VS is based on national legislation and has the flexibility and legal framework necessary in order to propose legislation and set regulations.

3. The VS is based on national legislation and proposes legislation and set regulations, applying procedures that take into consideration the opinions of its users.

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34 Regulations are sanitary measures that include all pertinent laws, decrees, regulations and technical prescriptions and procedures. Compliance is verified by VS through inspections and performance assessments

35 Regulations are sanitary measures that include all pertinent laws, decrees, regulations and technical prescriptions and procedures. Compliance is verified by VS through inspections and performance assessments
Appendix VI (contd)

3. **International harmonization**

The capability and authority of the VS to be active in international harmonization and ensure that the national laws and regulation covered under its mandate are in conformity with relevant international standards, guidelines and recommendations.

Levels of advancement:

0. The VS has no process to be aware of international standards. National laws and regulation do not take account of international standards, guidelines and recommendations.

1. The VS is aware of relevant standards but has no process to identify gaps, inconsistencies, or non-conformities in national laws and regulation as compared to international standards, guidelines and recommendations.

2. The VS monitors the establishment of new international standards, guidelines and recommendations and periodically reviews national laws and regulation with the aim of harmonizing them as appropriate with international standards, guidelines and recommendations.

3. Same as previous level plus the VS is active in reviewing and commenting on draft standards, guidelines and recommendations to relevant intergovernmental organizations.

4. Same as previous level plus the VS actively and regularly participates at the international level in the formulation, negotiation and adoption of international standards, guidelines and recommendations.  

4. **Certification**

The capability and authority of the VS to certify products, services and processes covered under its mandate and in accordance with the national laws and regulations and international standards, guidelines and recommendations.

Levels of advancement:

0. The VS has neither the capability nor the authority to certify animal health status, products, services or processes.

1. The VS has the authority to certify selected animals, animal products, services or processes.

2. The VS carries out certification programs for selected animals, animal products, services or processes.

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36 A country could be active in international standard setting without actively pursuing national changes. The importance of this element is to promote national change.

37 All certification procedures have to take into account the OIE standards on quality of VS and on certification.

In carrying out certification programmes, the VS must always operate free of political interference from the private sector. However some of these programmes can be executed by independent parties, which have been delegated and audited by the Veterinary Services.
3. The VS can develop and carry out certification programs for all animals, animal products, services or processes.

4. The veterinary service has certification power as necessary for all relevant animals and animal products and carries out audits of its certification programs through an efficient national chain of command in order to maintain confidence in its system.

5. **Equivalency** and other sanitary agreements

The capability and authority of the VS to negotiate implement and maintain equivalency and other sanitary agreements with other countries on veterinary requirements under its mandate.

Levels of advancement:

0. The VS has neither the authority nor the capability to negotiate and approve equivalency and other sanitary agreements with other countries.

1. The VS has the authority to negotiate and approve equivalency and other sanitary agreements with other countries.

2. Same as previous level plus the VS evaluates and proposes equivalency and other sanitary agreements with other countries on selected animals, animal products and processes.

3. Same as previous level plus the VS actively pursues the development of equivalency and other sanitary agreements with other countries on new products and processes.

4. Same as previous level plus the VS has a program that includes the feedback of its users along with advances in international standards, guidelines and recommendations, and then pursues specific equivalency and other sanitary agreements with other countries.

6. **Traceability**

The capability and authority of the VS to track the history, location and distribution of animals and their related products covered under its mandate.

Levels of advancement:

0. The VS has no program to track animals and their related products.

1. The VS can document and inspect the sanitary status at specific points across the agro-food chain for selected animals and their related products.

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38 The term, equivalency, refers to the state wherein the sanitary measure(s) proposed by the exporting country as an alternative to those of the importing country, achieve(s) the same level of protection. Guidelines on equivalency published in the OIE Codes have to be taken into account.

39 In compliance with OIE definitions, guidelines and relevant chapters of the Code on certain diseases.
Appendix VI (contd)

2. The VS has procedures in place and can track and inspect selected animals and their related products across that portion of the agri-food chain covered under its mandate.

3. The VS, along with the other relevant State institutions and its users, has coordinated procedures in place that can track and inspect animals and related animal products across the entire agri-food chain.

4. The VS, in cooperation with the other relevant State institutions and its users, carries out audits of its traceability procedures.

5. The VS manage and/or inspect a national data base on relevant animals and their movements.

7. Transparency

The capability and authority of the VS to notify the WTO/SPS and the OIE of its national regulations, sanitary status and decisions on the control of relevant diseases, in accordance with the obligations, standards and procedures established by these organizations.

Levels of advancement:

0. The VS does not notify the WTO/SPS and the OIE of its national regulations and decisions on control of relevant diseases, and the OIE of its sanitary status.

1. The VS partially notifies the WTO/SPS and the OIE of its national regulations and decisions on control of relevant diseases, and the OIE of its sanitary status.

2. The VS notifies the WTO/SPS and the OIE of its national regulations and decisions on control of relevant diseases, and the OIE of its sanitary status, in full compliance with the criteria established by these organizations.

3. The VS informs users of changes in its regulations and decisions on control of relevant diseases and sanitary status, changes in the regulations and sanitary status of other countries, and raises awareness with its users of the importance of being transparent.

4. The VS, along with the other relevant State institutions, carries out audits of its transparency procedures40 through an efficient national chain of command.

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40 In compliance with OIE standards on evaluation of VS
8. **Zoning**

The capability and authority of the VS to establish and maintain disease free zones or zones of low disease prevalence, in accordance to the criteria established by the WTO/SPS and the OIE.

Levels of advancement:

0. The VS cannot establish disease free zones or zones of low disease prevalence.

1. The national veterinary service can identify sub-populations to be regionalized, and establish the current sanitary status of selected animals and their related products originating from these prescribed areas.

2. The VS has implemented biosecurity control measures that enable it to establish disease free zones or zones of low disease prevalence for selected animals and their related products.

3. The VS collaborates with its users and relevant State institutions to define responsibilities execute actions and otherwise enable it to maintain disease free zones or zones of low disease prevalence for selected animals and their related products.

4. The VS demonstrates scientifically, the establishment of disease free zones/ or zones of low disease prevalence, and gains the recognition as such by other countries for selected animals and their related products.

5. The VS has a specific program that defines, establishes and demonstrates scientifically, new disease free zones or zones of low disease prevalence.

9. **Compartmentalization**

The capability and authority of the VS to establish and maintain disease free compartments or compartments of low disease prevalence, in accordance to the criteria established by the WTO/SPS and the OIE.

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41 For purposes of the Terrestrial Code and the OIE, ‘zoning’ and ‘regionalization’ have the same meaning. Implementation of these concepts has to take into account OIE standards included in the Codes.

42 The phrase, disease free zones: refers to animal sub-populations in which the absence of a given disease has been demonstrated to occur in accordance to the provisions outlined in the Terrestrial Animal Health Code of the OIE.

43 The phrase, zones of low disease prevalence, refers to zones, which can encompass the entire territory of a country, part of a country, or sub-populations within a country, in which a given disease exists only to a limited extent, and is subject to effective surveillance, control or eradication measures.

44 Implementation of this concepts has to take into account OIE standards included in the Codes.

45 The phrase, disease free compartments, refers to animal sub-populations in which the absence of a given disease has been demonstrated to occur in accordance to the provisions outlined in the Terrestrial Animal Health Code of the OIE.

46 The phrase, compartments of low disease prevalence, refers to compartments, which can encompass subpopulation within a compartment, in which a given disease exists only to a limited extent, and is subject to effective surveillance, control or eradication measures.
Appendix VI (contd)

Levels of advancement:

0. The VS cannot establish disease free compartments or compartments of low disease prevalence.

1. The national veterinary service can identify sub-populations to be regionalized, and establish the current sanitary status of selected animals and their related products originating from these prescribed areas.

2. The VS has implemented biosecurity control measures that enable it to establish disease free compartments or compartments of low disease prevalence for selected animals and their related products.

3. The VS collaborates with its users and relevant State institutions to define responsibilities execute actions and otherwise enable it to maintain disease free compartments or compartments of low disease prevalence for selected animals and their related products.

4. The VS demonstrates scientifically, the establishment of disease free compartments or compartments of low disease prevalence, and gains the recognition as such by other countries for selected animals and their related products.

5. The VS has a specific program that defines, establishes and demonstrates scientifically, new disease free compartments or compartments of low disease prevalence.
CHAPTER 1.3.5.

ZONING AND COMPARTMENTALISATION

Article 1.3.5.1.

Introduction

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

Given the difficulty of establishing and maintaining a disease free status for an entire country, especially for diseases the entry of which is difficult to control through measures at national boundaries, there may be benefits to Member Countries in establishing and maintaining a subpopulation with a different animal health status within national boundaries. Subpopulations may be separated by natural or artificial geographical barriers, or in certain animal industries, by the application of appropriate management systems, including biosecurity management.

Zoning and compartmentalisation are procedures implemented by a country under the provisions of this Chapter with a view to defining subpopulations of different animal health status within its territory for the purpose of disease control and/or international trade. Compartmentalisation applies to a subpopulation when management systems related to biosecurity are applied, while zoning applies when a subpopulation is defined on a geographical basis.

This chapter is to assist OIE Member Countries to establish and maintain different subpopulations within their national boundaries using the procedures of compartmentalisation and zoning. It also outlines a process for trading partners to follow in achieving recognition of such subpopulation. These procedures are 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 boundaries and within its territory.

The benefits of zoning and compartmentalisation may include a contribution to disease control or eradication within Member Countries, and to the safety of international trade. Zoning may encourage the more efficient use of resources within certain parts of a country to allow trade in certain commodities from that zone in accordance with this Terrestrial Code. Compartmentalisation may allow safe trade due to the functional separation of a sub-population from other domestic or wild animals through biosecurity measures, which a zone (through geographical separation alone) would not achieve. Following a disease outbreak, compartmentalisation may be able to take advantage of epidemiological linkages despite diverse geographical locations, to facilitate disease control.

Separate requirements will be developed for each disease for which the application of zoning or compartmentalisation is considered appropriate.
Appendix VII (contd)

Article 1.3.5.2.

General considerations

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 boundaries and within its territory.

The benefits of zoning and compartmentalisation may include a contribution to disease control or eradication within Member Countries, and to the safety of international trade. Zoning may encourage the more efficient use of resources within certain parts of a country to allow trade in certain commodities from that zone in accordance with this Terrestrial Code. Compartmentalisation may allow safe trade due to the functional separation of a sub-population from other domestic or wild animals through biosecurity measures, which a zone (through geographical separation alone) would not achieve. Following a disease outbreak, compartmentalisation may be able to take advantage of epidemiological linkages despite diverse geographical locations, to facilitate disease control.

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 measures stipulated in the relevant Chapters in this Terrestrial Code and should be able to explain to the Veterinary Services of an importing country the basis for its claim of a distinct animal health status for the zone or compartment in such terms.

The procedures used to establish and maintain the distinct 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, applicable biosecurity measures (including movement controls, use of natural and artificial boundaries, commercial management and husbandry practices), and surveillance and monitoring. The exporting country should be able to demonstrate, through detailed documentation published through official channels, that it has implemented the measures stipulated in this 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 Veterinary Administration of the exporting country certifies that the appropriate measures recommended in this Terrestrial Code are applied.

Article 1.3.5.3.

Prerequisite considerations in defining a zone or compartment

The exporting country should conduct a practical 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).
Article 1.3.5.4.

Principles for defining a zone or compartment

In conjunction with the above considerations, defining a zone or compartment should be based on the application of the following principles:

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

2. The requirements regarding a compartment should be established by the Veterinary Administration on the basis of relevant criteria such as biosecurity management and husbandry practices, and made public through official channels.

3. Animals and herds belonging to subpopulations need to be clearly recognizable as such. The Veterinary Administration must document in detail the measures taken to ensure the identification of the subpopulation and the recognition and maintenance of its health status.

4. The requirements necessary to preserve the distinct health status of a zone or compartment must be appropriate to the particular disease and will depend on the epidemiology of the disease, environmental factors, biosecurity management, animal husbandry practices, control measures and surveillance.

5. Thus defined, the zones and compartments constitute the relevant subpopulations for the application of the recommendations in Part 2 of this Terrestrial Code.

Article 1.3.5.5.

Sequence of steps to be taken in defining a zone/compartment

There is no single sequence of steps which must be followed in defining a zone or a compartment. The steps that the Veterinary Services of importing and exporting countries choose and implement will generally depend on the circumstances existing within a country and at its borders. The recommended steps are:

1. For zoning
   a) The exporting country identifies a geographical area within its territory which it considers to contain an animal subpopulation with a distinct health status with respect to a specific disease/specific diseases, based on surveillance and monitoring.
   b) The exporting country identifies the procedures which are being, or could be, employed to distinguish such an area epidemiologically from other parts of its territory, in accordance with the measures stipulated in this Terrestrial Code.
   c) The exporting country provides the information above to the importing country, and explains that the area can be treated as an epidemiologically separated zone for international trade purposes.
Appendix VII (contd)

d) The importing country determines whether it may accept such an area as a zone for the importation of animals and animal products, taking into account:

i) an evaluation of the exporting country's Veterinary Services;

ii) the result of a risk assessment based on the information provided by the exporting country and its own research;

iii) its own animal health situation with respect to the disease(s) concerned; and

iv) other relevant OIE standards.

e) The importing country notifies the exporting country of the result of its determination and the underlying reasons, within a reasonable period of time, being either:

i) recognition of the zone;

ii) request for further information; or

iii) rejection of the area as a zone for international trade purposes.

f) An attempt should be made to resolve any differences of opinion over the definition of the zone, either in the interim or finally, by using an agreed mechanism to reach consensus (such as the OIE dispute settlement mechanism).

g) The importing country and the exporting country may enter into a formal agreement defining the zone.

2. For compartmentalisation

a) Based on discussions with the relevant enterprise/industry, the Veterinary Administration of the exporting country identifies within its territory one or more establishments or other premises owned by an enterprise(s) which operates under a common biosecurity management system, and which it considers contains an identifiable animal subpopulation with a distinct health status with respect to a specific disease/specific diseases, and that this status is maintained through a partnership between the relevant enterprise/industry and the Veterinary Services of the exporting country.

b) The exporting country examines the ‘biosecurity management manual’ produced by the enterprise/industry for such establishment(s), and confirms through an audit that:

i) such establishment(s) is(are) epidemiologically closed throughout its routine operating procedures as a result of effective implementation of its ‘biosecurity management manual’ and;

ii) the surveillance and monitoring programme in place is appropriate to verify the free status of such establishment(s) with respect to such disease(s).

c) The exporting country identifies such an enterprise to be a free compartment, in accordance with the measures stipulated in this Terrestrial Code.

d) The exporting country provides the information above to the importing country, and explains that such an enterprise can be treated as an epidemiologically separated compartment for international trade purposes.
Appendix VII (contd)

e) The importing country determines whether it may accept such an enterprise as a compartment taking into account:
   i) an evaluation of the exporting country’s Veterinary Services;
   ii) the result of a risk assessment based on the information provided by the exporting country and its own research;
   iii) its own animal health situation with respect to the disease(s) concerned; and
   iv) other relevant OIE standards.

f) The importing country notifies the exporting country of the result of its examination and the underlying reasons, within a reasonable period of time, being either:
   i) recognition of the compartment;
   ii) request for further information; or
   iii) rejection of such an enterprise as a compartment for international trade purposes.

g) An attempt should be made to resolve any differences of opinion over the definition of the compartment, either in the interim or finally, by using an agreed mechanism to reach consensus (such as the OIE dispute settlement mechanism).

h) The importing country and the exporting country may enter into a formal agreement defining the compartment.
CHAPTER 2.1.1.
CRITERIA FOR LISTING DISEASES

Article 2.1.1.1.

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

<table>
<thead>
<tr>
<th>Basic criteria</th>
<th>Parameters (at least one ‘yes’ answer means that the criterion has been met)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>International Spread</strong></td>
<td>Has international spread been proven on three or more occasions? <strong>OR</strong></td>
</tr>
<tr>
<td></td>
<td>Are more than three countries with populations of susceptible animals free of the disease or facing impending freedom (based on the Terrestrial Code provisions, especially Appendix 3.8.1)? <strong>OR</strong></td>
</tr>
<tr>
<td></td>
<td>Do OIE annual reports indicate that a significant number of countries with susceptible populations have reported absence of the disease for several consecutive years?</td>
</tr>
<tr>
<td><strong>Zoonotic Potential</strong></td>
<td>Has transmission to humans been proven? (with the exception of artificial circumstances) <strong>AND</strong></td>
</tr>
<tr>
<td></td>
<td>Is human infection associated with severe consequences? (death or prolonged illness)</td>
</tr>
<tr>
<td><strong>Significant Spread within Naïve Populations</strong></td>
<td>Does the disease exhibit significant mortality at the level of a country or zone/compartment? <strong>AND/OR</strong></td>
</tr>
<tr>
<td></td>
<td>Does the disease exhibit significant morbidity at the level of a country or zone/compartment?</td>
</tr>
<tr>
<td><strong>Emerging Diseases</strong></td>
<td>Is there rapid spread and/or apparent zoonotic properties?</td>
</tr>
</tbody>
</table>
Appendix VIII (contd)

Article 2.1.1.2.

The criteria in Article 2.1.1. above are applied according to the decision-making model shown below:

![Decision Tree Diagram]

**INTERNATIONAL SPREAD**
- Has international spread been proven on three or more occasions? **OR**
- Are more than three countries with populations of susceptible animals free of the disease or facing impending freedom (based on Code provisions, especially Appendix 3.8.1)? **OR**
- Do OIE annual reports indicate that a significant number of countries with susceptible populations have reported absence of the disease for several consecutive years?

**EMERGING**
- (A newly recognised pathogen or known pathogen behaving differently)
- Is there rapid spread or apparent zoonotic properties?

**ZOONOTIC**
- Has transmission to humans been proven? (with the exception of artificial circumstances) **AND**
- Is human infection associated with severe consequences? (death or prolonged illness)

**SIGNIFICANT SPREAD IN NAIVE POPULATIONS**
- Does the disease exhibit significant mortality at the level of a country or zone? **OR**
- Does the disease exhibit significant morbidity at the level of a country or zone?

**Article 2.1.1.3.**

The following diseases are included in the OIE List.

1. The following diseases are included within the category of multiple species diseases:
   - Anthrax
   - Aujeszky's disease
   - Bluetongue
   - Brucellosis (*Brucella abortus*)
   - Brucellosis (*Brucella melitensis*)
- Brucellosis (*Brucella suis*)
- Crimean Congo haemorrhagic fever
- Echinococcosis/hydatidosis
- Foot and mouth disease
- Heartwater
- Japanese encephalitis
- Leptospirosis
- New world screwworm (*Cochliomyia hominivorax*)
- Old world screwworm (*Chrysomya bezziana*)
- Paratuberculosis
- Q fever
- Rabies
- Rift Valley fever
- Rinderpest
- Trichinellois
- Tularemia
- Vesicular stomatitis
- West Nile fever.

2. The following diseases are included within the category of cattle diseases:
- Bovine anaplasmosis
- Bovine babesiosis
- Bovine genital campylobacteriosis
- Bovine spongiform encephalopathy
- Bovine tuberculosis
- Bovine viral diarrhoea
Appendix VIII (contd)

- Contagious bovine pleuropneumonia.
- Enzootic bovine leukosis
- Haemorrhagic septicaemia
- Infectious bovine rhinotracheitis/infectious pustular vulvovaginitis
- Lumpy skin disease
- Malignant catarrhal fever
- Theileriosis
- Trichomonosis
- Trypanosomosis (tsetse-transmitted).

3. The following diseases are included within the category of sheep and goat diseases:

- Caprine arthritis/encephalitis
- Contagious agalactia
- Contagious caprine pleuropneumonia
- Enzootic abortion of ewes (ovine chlamydiosis)
- Maedi–visna
- Nairobi sheep disease
- Ovine epididymitis (*Brucella ovis*)
- Peste des petits ruminants
- Salmonellosis (*S. abortusovis*)
- Scrapie
- Sheep pox and goat pox.

4. The following diseases are included within the category of equine diseases:

- African horse sickness
- Contagious equine metritis
- Dourine
- Equine encephalomyelitis (Eastern)
- Equine encephalomyelitis (Western)
- Equine infectious anaemia
- Equine influenza
- Equine piroplasmosis
- Equine rhinopneumonitis
- Equine viral arteritis
- Glanders
- Surra (*Trypanosoma evansi*)
- Venezuelan equine encephalomyelitis.

5. The following diseases are included within the category of swine diseases:
   - African swine fever
   - Classical swine fever
   - Nipah virus encephalitis
   - Porcine cysticercosis
   - Porcine reproductive and respiratory syndrome
   - Swine vesicular disease
   - Transmissible gastroenteritis.

6. The following diseases are included within the category of avian diseases:
   - Avian chlamydiosis
   - Avian infectious bronchitis
   - Avian infectious laryngotracheitis
   - Avian mycoplasmosis (*M. gallisepticum*)
   - Avian mycoplasmosis (*M. synoviae*)
   - Duck virus hepatitis
Appendix VIII (contd)

- Fowl cholera
- Fowl typhoid
- Highly pathogenic avian influenza
- Infectious bursal disease (Gumboro disease)
- Marek's disease
- Newcastle disease
- Pullorum disease
- Turkey rhinotracheitis.

7. The following diseases are included within the category of lagomorph diseases:
   - Myxomatosis
   - Rabbit haemorrhagic disease.

8. The following diseases are included within the category of bee diseases:
   - Acarapisosis of honey bees
   - American foulbrood of honey bees
   - European foulbrood of honey bees
   - Small hive beetle infestation (Aethina tumida)
   - Tropilaelaps infestation of honey bees
   - Varroosis of honey bees.

9. The following diseases are included within the category of other diseases:
   - Camelpox
   - Leishmaniosis.

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CHAPTER 2.2.10.

FOOT AND MOUTH DISEASE

Article 2.2.10.1.

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.

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 RNA specific to one or more of the serotypes of FMDV has been identified in samples from 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, 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 2.2.10.2.

FMD free country where vaccination is not practised

To qualify for inclusion in the existing list of FMD free countries where vaccination is not practised, a country 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,
   c) no vaccination against FMD has been carried out during the past 12 months,

   and supply documented evidence that surveillance for both FMD and FMDV infection in accordance with Appendix 3.8.7. is in operation and that regulatory measures for the prevention and control of FMD have been implemented;
Appendix IX (contd)

3. not have imported since the cessation of vaccination any animals vaccinated against FMD.

The country will be included in the list only after the submitted evidence has been accepted by the OIE.

Article 2.2.10.3.

FMD free country where vaccination is practised

To qualify for inclusion in the list of FMD free countries where vaccination is practised, a country 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 Appendix 3.8.7. 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 country will be included in the list only after the submitted evidence has been accepted by the OIE.

If an FMD free country where vaccination is practised wishes to change its status to FMD free country where vaccination is not practised, the country should wait for 12 months after vaccination has ceased and provide evidence showing that FMDV circulation has not occurred during that period.

Article 2.2.10.4.

FMD free zone where vaccination is not practised

An FMD free zone where vaccination is not practised can be established in either an FMD free country where vaccination is practised or in a country of which parts are infected. Susceptible animals in the FMD free zone should be separated from the rest of the country, if infected, and from neighbouring infected countries by a buffer zone, or physical or geographical barriers, and animal health measures that effectively prevent the entry of the virus should be implemented. A country in which an FMD free zone where vaccination is not practised is to be established 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 an 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 Articles 2.2.10.8.;
3. supply documented evidence that surveillance for both FMD and FMDV infection in accordance with Appendix 3.8.7. is in operation in the proposed FMD free zone where vaccination is not practised;

4. describe in detail:
   a) regulatory measures for the prevention and control of both FMD and FMDV infection,
   b) the boundaries of the FMD free zone and, if applicable, the buffer 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 FMDV free zone (in particular if the procedure described in Article 2.2.10.8. 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.

Article 2.2.10.5.

FMD free zone where vaccination is practised

An FMD free zone where vaccination is practised can be established in either an FMD free country where vaccination is not practised or in a country of which parts are infected. Susceptible animals in the FMD free zone where vaccination is practised should be separated from the rest of the country, if infected, and from neighbouring infected countries by a buffer zone, or physical or geographical barriers. Animal health measures that effectively prevent the entry of the virus should be implemented.

Vaccination of zoo animals, animals belonging to rare species or breeds, or animals in research centres as a precaution for conservation purposes is an example of implementation of an FMD free zone or compartment where vaccination is practised.

A country in which an FMD free zone where vaccination is practised is to be established should:

1. have a record of regular and prompt animal disease reporting;

2. send a declaration to the OIE that it wishes to establish an FMD free zone where vaccination is practised, where 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 surveillance for FMD and FMDV circulation in accordance with Appendix 3.8.7. is in operation in the proposed FMD free zone;

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,
Appendix IX (contd)

b) the boundaries of the FMD free zone where vaccination is practised and, if applicable, the buffer zone or physical or geographical barriers.

c) the system for preventing the entry of the virus into the FMD free zone (in particular if the procedure described in Article 2.2.10.8. is implemented),

and supply evidence that these are properly implemented and supervised;

5. supply documented evidence that it has a system of intensive and frequent surveillance for FMD and FMDV circulation in the FMD free zone where vaccination is practised.

The 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.

If a country that has an FMD free zone where vaccination is practised wishes to change the status of the zone to FMD free zone where vaccination is not practised, a waiting period of 12 months after vaccination has ceased is required and evidence must be provided showing that FMDV infection has not occurred in the said zone during that period.

Article 2.2.10.6.

FMD infected country or zone

An FMD infected country is a country that does not fulfil the requirements to qualify as either an FMD free country where vaccination is not practised or an FMD free country where vaccination is practised.

An FMD infected zone is a zone that does not fulfil the requirements to qualify as either an FMD free zone where vaccination is not practised or an FMD free zone where vaccination is practised.

Article 2.2.10.7.

Recovery of free status

1. When an FMD outbreak or FMDV infection occurs in an 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 Appendix 3.8.7.; 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 Appendix 3.8.7.; 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 Appendix 3.8.7., 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 2.2.10.2 or 2.2.10.4. applies.
2. When an FMD outbreak or FMDV infection occurs in an 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 Appendix 3.8.7. 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 Appendix 3.8.7. 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 2.2.10.8.

Transfer directly to slaughter of FMD susceptible animals from an infected zone to a free zone 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 zone directly to slaughter.

In the absence of an abattoir in the buffer 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 Appendix 3.6.2.

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 2.2.10.11.
Appendix IX (contd)

Article 2.2.10.9.
When importing from FMD free countries or zones where vaccination is not practised, Veterinary Administrations should require:

for FMD susceptible animals
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 an FMD free country or zone where vaccination is not practised since birth or for at least the past 3 months.

Article 2.2.10.10.
When importing from FMD free countries or zones where vaccination is practised, Veterinary Administrations should require:

for domestic ruminants and pigs
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 an FMD free country 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 an FMD free country or zone where vaccination is not practised.

Article 2.2.10.11.
When importing from FMD infected countries or zones, Veterinary Administrations should require:

for domestic ruminants and pigs
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 10-kilometre radius of the establishment of origin for the relevant period as defined in points a) and b) above; and
Appendix IX (contd)

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 10-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 10-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 2.2.10.12.

When importing from FMD free countries or zones where vaccination is not practised, Veterinary Administrations should require:

for fresh semen of domestic ruminants and pigs

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 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 Appendix 3.2.1. or Appendix 3.2.2., as relevant.

Article 2.2.10.13.

When importing from FMD free countries or zones where vaccination is not practised, Veterinary Administrations should require:

for frozen semen of domestic ruminants and pigs

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 an 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 Appendix 3.2.1. or Appendix 3.2.2., as relevant.
Appendix IX (contd)

Article 2.2.10.14.

When importing from FMD free countries or zones where vaccination is practised, Veterinary Administrations should require:

for semen of domestic ruminants and pigs

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 country or zone free from FMD for at least 3 months prior to collection;
   c) if destined to an 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 Appendix 3.2.1. or Appendix 3.2.2., 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 2.2.10.15.

When importing from FMD infected countries or zones, Veterinary Administrations should require:

for semen of domestic ruminants and pigs

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;
Appendix IX (contd)

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 Appendix 3.2.1. or Appendix 3.2.2., 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 2.2.10.16.

Irrespective of the FMD status of the exporting country or zone, Veterinary Administrations 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 Appendix 3.3.1. or Appendix 3.3.3., as relevant.

Article 2.2.10.17.

When importing from FMD free countries or zones where vaccination is not practised, Veterinary Administrations should require:

for in vitro produced embryos of cattle

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 in a country or zone free from FMD at the time of collection;

2. fertilisation was achieved with semen meeting the conditions referred to in Articles 2.2.10.12., 2.2.10.13., 2.2.10.14. or 2.2.10.15., as relevant;

3. the oocytes were collected, and the embryos were processed and stored in conformity with the provisions of Appendix 3.3.2. or Appendix 3.3.3., as relevant.
Appendix IX (contd)

Article 2.2.10.18.

When importing from FMD free countries or zones where vaccination is practised, Veterinary Administrations should require:

_for in vitro produced embryos of cattle_

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 in a country or zone free from FMD for at least 3 months prior to collection;
   c) if destined for an 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 2.2.10.12., 2.2.10.13., 2.2.10.14. or 2.2.10.15., as relevant;

4. the oocytes were collected, and the embryos were processed and stored in conformity with the provisions of Appendix 3.3.2. or Appendix 3.3.3., as relevant.

Article 2.2.10.19.

When importing from FMD free countries or zones where vaccination is not practised, Veterinary Administrations should require:

_for fresh meat of FMD susceptible animals_

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 2.2.10.9., Article 2.2.10.10. or Article 2.2.10.11.;

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 2.2.10.20.

When importing from FMD free countries where vaccination is practised or from FMD free zones where vaccination is practised, Veterinary Administrations should require:
Appendix IX (contd)

for fresh meat of cattle and buffalo (Bubalus bubalis) (excluding feet, head and viscera)

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 2.2.10.9., Article 2.2.10.10. or Article 2.2.10.11.;

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 2.2.10.21.

When importing from FMD free countries where vaccination is practised or from FMD free zones where vaccination is practised, Veterinary Administrations should require:

for fresh meat or meat products of pigs and ruminants other than cattle and buffalo

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 2.2.10.9., Article 2.2.10.10. or Article 2.2.10.11.;

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 2.2.10.22.

When importing from FMD infected countries or zones, where an official control programme exists, involving compulsory systematic vaccination of cattle, Veterinary Administrations should require:

for fresh meat of cattle and buffalo (Bubalus bubalis) (excluding feet, head and viscera)

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 10-kilometre radius of the establishment during that period;
Appendix IX (contd)

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 lymph 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.

Article 2.2.10.23.

When importing from FMD infected countries or zones, Veterinary Administrations should require:

for meat products of domestic ruminants and pigs

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 3.6.2.1.;

3. the necessary precautions were taken after processing to avoid contact of the meat products with any potential source of FMD virus.

Article 2.2.10.24.

When importing from FMD free countries or zones (where vaccination either is or is not practised), Veterinary Administrations should require:

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

the presentation of an international veterinary certificate attesting that these products come from animals which have been kept in the country or zone since birth, or which have been imported in accordance with Article 2.2.10.9., Article 2.2.10.10. or Article 2.2.10.11.
Article 2.2.10.25.

When importing from FMD infected countries or zones where an official control programme exists, Veterinary Administrations should require:

_for milk, cream, milk powder and milk products_

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 3.6.2.5. and in Article 3.6.2.6.;

2. the necessary precautions were taken after processing to avoid contact of the products with any potential source of FMD virus.

Article 2.2.10.26.

When importing from FMD infected countries, Veterinary Administrations should require:

_for blood and meat-meals (from domestic or wild ruminants and pigs)_

the presentation of an international veterinary certificate attesting that the manufacturing method for these products included heating to a minimum core internal temperature of 70°C for at least 30 minutes.

Article 2.2.10.27.

When importing from FMD infected countries, Veterinary Administrations should require:

_for wool, hair, bristles, raw hides and skins (from domestic or wild ruminants and pigs)_

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 3.6.2.2., 3.6.2.3. and 3.6.2.4.;

2. the necessary precautions were taken after collection or processing to avoid contact of the products with any potential source of FMD virus.

Veterinary Administrations 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.
Appendix IX (contd)

Article 2.2.10.28.
When importing from FMD infected countries or zones, Veterinary Administrations should require:
for straw and forage
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 2.2.10.29.
When importing from FMD free countries or zones (where vaccination either is or is not practised), Veterinary Administrations should require:
for skins and trophies derived from FMD susceptible wild animals
the presentation of an international veterinary certificate attesting that these products are derived from animals that have been kept in such a country or zone since birth, or which have been imported from a country or zone free of FMD (where vaccination either is or is not practised).

Article 2.2.10.30.
When importing from FMD infected countries or zones, Veterinary Administrations should require:
for skins and trophies derived from FMD susceptible wild animals
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 3.6.2.7.

[Note: International veterinary certificates for animal products coming from infected countries or zones may not be required if the products are transported in an approved manner to premises controlled and approved by the Veterinary Administration of the importing country for processing to ensure the destruction of the FMD virus in conformity with the procedures referred to in Articles 3.6.2.2., 3.6.2.3. and 3.6.2.4.]
APPENDIX 3.8.7.

GUIDELINES FOR THE SURVEILLANCE OF FOOT AND MOUTH DISEASE

Article 3.8.7.1.

Introduction

This Appendix defines the principles and provides a guide for the surveillance of foot and mouth disease (FMD) in accordance with Appendix 3.8.1, applicable to countries seeking recognition from the OIE for freedom from FMD, either with or without the use of vaccination. This may be for the entire country or a zone or compartment within the country. Guidance for countries seeking reestablishment of freedom from FMD for the whole country or a zone or compartment, either with or without vaccination, following an outbreak, as well as guidelines for the maintenance of FMD status are provided. These guidelines are intended to expand on and explain the requirements of Chapter 2.2.10. 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 guidelines 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 applicant country 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 Member Countries 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 Appendix, virus circulation means transmission of FMDV as demonstrated by clinical signs, serological evidence or virus isolation.

Article 3.8.7.2.

General conditions and methods

1. A surveillance system in accordance with Appendix 3.8.1 should be under the responsibility of the Veterinary Administration. 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.
Appendix X (contd)

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 Administration. 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 an approved 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 an FMD infected country or 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 standstill orders, etc.).

Article 3.8.7.3.

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 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 applicant country should justify the surveillance strategy chosen as adequate to detect the presence of FMDV infection/circulation in accordance with Appendix 3.8.1. 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 Country wishes to apply for recognition of a specific zone or compartment 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 or compartment.

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 applicant country must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Appendix 3.8.1. 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.
Appendix X (contd)

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);

dl) heterophile (cross) reactions.

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 Appendix 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 3.8.7.4.

Countries applying for freedom from FMD for the whole country or a zone or a compartment where vaccination is not practised

In addition to the general conditions described in Chapter 2.2.10., a Member Country applying for recognition of FMD freedom for the country or a zone or a compartment 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 Appendix, 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.
Appendix X (contd)

Article 3.8.7.5.

Countries, or zones or compartments applying for freedom from FMD where vaccination is practised

In addition to the general conditions described in Chapter 2.2.10., a Member Country applying for recognition of country or zone or compartment freedom from FMD with vaccination should show evidence of an effective surveillance programme planned and implemented according to general conditions and methods in this Appendix. Absence of clinical disease in the country, or zone or compartment 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 or compartment, 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 3.8.7.6.

Countries, or zones or compartments re-applying for freedom from FMD where vaccination is either practised or not practised, following an outbreak

In addition to the general conditions described in Chapter 2.2.10., a country re-applying for country, or zone or compartment 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 zone or compartment 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 re-instatement of freedom from FMD depends on which of these alternatives is followed. The time periods are prescribed in Article 2.2.10.7.

In all circumstances, a Member Country re-applying for country, or zone or compartment 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 Appendix.

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Appendix X (contd)

Article 3.8.7.7.

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.

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;
Appendix X (contd)

– 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.
Figure 1 Schematic representation of laboratory tests for determining evidence of FMDV infection through or following serological surveys

Key:

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ELISA</td>
<td>Enzyme-linked immunosorbent assay</td>
</tr>
<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>
</tr>
<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>OP</td>
<td>Oesophageal pharyngeal sample</td>
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<tr>
<td>SP</td>
<td>Structural protein test</td>
</tr>
<tr>
<td>S</td>
<td>No evidence of FMDV</td>
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</tbody>
</table>
CHAPTER 2.2.13. BLUETONGUE

Article 2.2.13.1.

For the purposes of the Terrestrial Code, the infective period for bluetongue virus (BTV) shall be 60 days.

The global BTV distribution is currently between latitudes of approximately 50°N and 34°S but is known to be expanding in the northern hemisphere.

In the absence of clinical disease in a country or zone within this part of the world, its BTV status should be determined by an ongoing surveillance and monitoring programme (in accordance with Appendix 3.8.X.) designed in accordance with the epidemiology of the disease, i.e. focusing on climatic and geographical factors, the biology and likely competence of Culicoides and/or serology of susceptible animals. The programme may need to be adapted to target parts of the country or zone at a higher risk due to historical, geographical and climatic factors, ruminant population data and Culicoides ecology, or proximity to enzootic or incursional zones as described in Appendix 3.8.X.

All countries or zones adjacent to a country or zone not having free status should be subjected to similar surveillance. 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 BTV, or a bluetongue surveillance programme (in accordance with Appendix 3.8.X.) in the country or zone not having free status supports a lesser distance.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 2.2.13.2.

BTV free country or zone

1. A country or a zone may be considered free from BTV when bluetongue is notifiable in the whole country and either:
   a) the country or zone lies wholly north of 50°N or south of 34°S, and is not adjacent to a country or zone not having a free status; or
   b) a surveillance and monitoring programme in accordance with Appendix 3.8.X. has demonstrated no evidence of BTV in the country or zone during the past 2 years; or
   c) a surveillance and monitoring programme has demonstrated no evidence of Culicoides likely to be competent BTV vectors in the country or zone.

2. A BTV free country or zone in which surveillance and monitoring has found no evidence that Culicoides likely to be competent BTV vectors are present will not lose its free status through the importation of vaccinated, seropositive or infective animals, or semen or embryos/ova from infected countries or zones.
Appendix XI (contd)

3. A BTV free country or zone in which surveillance and monitoring has found evidence that *Culicoides* likely to be competent BTV vectors are present will not lose its free status through the importation of vaccinated or seropositive animals from infected countries or zones, provided:

   a) the animals have been vaccinated in accordance with the *Terrestrial Manual* at least 60 days prior to dispatch with a vaccine which covers all serotypes whose presence in the source population has been demonstrated through a surveillance and monitoring programme in accordance with Appendix 3.8.X., and that the animals are identified in the accompanying certification as having been vaccinated; or

   b) the animals are not vaccinated, and a surveillance and monitoring programme in accordance with Appendix 3.8.X. has been in place in the source population for a period of 60 days immediately prior to dispatch, and no evidence of BTV transmission has been detected.

4. A BTV free country or zone adjacent to an infected country or zone should include a zone in which surveillance is conducted in accordance with Appendix 3.8.X. Animals within this zone must be subjected to continuing surveillance. The boundaries of this zone must be clearly defined, and must take account of geographical and epidemiological factors that are relevant to BTV transmission.

   Article 2.2.13.3.

**BTV seasonally free zone**

A BTV seasonally free zone is a part of an infected country or zone for which for part of a year, surveillance and monitoring demonstrate no evidence either of BTV transmission or of adult *Culicoides* likely to be competent BTV vectors.

For the application of Articles 2.2.13.7., 2.2.13.10. and 2.2.13.14., the seasonally free period is taken to commence the day following the last evidence of BTV transmission (as demonstrated by the surveillance and monitoring programme), or of the cessation of activity of adult *Culicoides* likely to be competent BTV vectors.

For the application of Articles 2.2.13.7., 2.2.13.10. and 2.2.13.14., the seasonally free period is taken to conclude either:

1. at least 28 days before the earliest date that historical data show bluetongue virus activity has recommenced; or

2. immediately if current climatic data or data from a surveillance and monitoring programme indicate an earlier resurgence of activity of adult *Culicoides* likely to be competent BTV vectors.

A BTV seasonally free zone in which surveillance and monitoring has found no evidence that *Culicoides* likely to be competent BTV vectors are present will not lose its free status through the importation of vaccinated, seropositive or infective animals, or semen or embryos/ova from infected countries or zones.

   Article 2.2.13.4.

**BTV infected country or zone**

A BTV infected country or zone is a clearly defined area where evidence of BTV has been reported during the past 2 years.
Article 2.2.13.5.

Veterinary Administrations of countries shall consider whether there is a risk with regard to BTV infection in accepting importation or transit through their territory, from other countries, of the following commodities:

1. ruminants and other BTV susceptible herbivores;
2. semen of these species;
3. embryos/ova of these species;
4. pathological material and biological products (from these species) (see Chapter 1.4.6. and Section 1.5.).

Other commodities should be considered as not having the potential to spread BTV when they are the subject of international trade.

Article 2.2.13.6.

When importing from BTV free countries or zones, Veterinary Administrations should require:

for ruminants and other BTV susceptible herbivores

the presentation of an international veterinary certificate attesting that:

1. the animals were kept in a BTV free country or zone since birth or for at least 60 days prior to shipment; or
2. the animals were kept in a BTV free country or zone for at least 28 days, then were subjected, with negative results, to a serological test to detect antibody to the BTV group according to the Terrestrial Manual and remained in the BTV free country or zone until shipment; or
3. the animals were kept in a BTV free country or zone for at least 7 days, then were subjected, with negative results, to an agent identification test according to the Terrestrial Manual and remained in the BTV free country or zone until shipment; or
4. the animals:
   a) were kept in a BTV free country or zone for at least 7 days;
   b) were vaccinated in accordance with the Terrestrial Manual 60 days before introduction into the free country or zone against all serotypes whose presence in the source population has been demonstrated through a surveillance and monitoring programme as described in Appendix 3.8.1.;
   c) were identified as having been vaccinated; and
   d) remained in the BTV free country or zone until shipment;
Appendix XI (contd)

AND

5. 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 attack from Culicoides likely to be competent BTV vectors at all times when transiting through an infected zone; or
   c) had been vaccinated in accordance with point 4) above.

Article 2.2.13.7.

When importing from BTV seasonally free zones, Veterinary Administrations should require:

for ruminants and other BTV susceptible herbivores

the presentation of an international veterinary certificate attesting that the animals:

1. were kept during the seasonally free period in a BTV seasonally free zone for at least 60 days prior to shipment; or

2. were kept during the BTV seasonally free period in a BTV seasonally free zone for at least 28 days prior to shipment, and were subjected during the residence period in the zone to a serological test to detect antibody to the BTV group according to the Terrestrial Manual, with negative results, carried out at least 28 days after the commencement of the residence period; or

3. were kept during the BTV seasonally free period in a BTV seasonally free zone for at least 14 days prior to shipment, and were subjected during the residence period in the zone to an agent identification test according to the Terrestrial Manual, with negative results, carried out at least 14 days after the commencement of the residence period; or

4. were kept during the seasonally free period in a BTV seasonally free zone, and were vaccinated in accordance with the Terrestrial Manual 60 days before introduction into the free country or zone against all serotypes whose presence in the source population has been demonstrated through a surveillance and monitoring programme in accordance with Appendix 3.8.X., were identified as having been vaccinated and remained in the BTV free country or zone until shipment;

AND

5. 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 attack from Culicoides likely to be competent BTV vectors at all times when transiting through an infected zone; or
   c) were vaccinated in accordance with point 4) above.
Article 2.2.13.8.

When importing from BTV infected countries or zones, Veterinary Administrations should require:

the presentation of an international veterinary certificate attesting that the animals:

1. were protected from attack from Culicoides likely to be competent BTV vectors for at least 60 days prior to shipment; or

2. were protected from attack from Culicoides likely to be competent BTV vectors for at least 28 days prior to shipment, and were subjected during that period to a serological test according to the Terrestrial Manual to detect antibody to the BTV group, with negative results, carried out at least 28 days after introduction into the quarantine station; or

3. were protected from attack from Culicoides likely to be competent BTV vectors for at least 14 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 at least 14 days after introduction into the quarantine station; or

4. were vaccinated in accordance with the Terrestrial Manual at least 60 days before shipment, against all serotypes whose presence in the source population has been demonstrated through a surveillance and monitoring programme in accordance with Appendix 3.8.1., and were identified in the accompanying certification as having been vaccinated; or

5. are not vaccinated, a surveillance and monitoring programme in accordance with 3.8.1. has been in place in the source population for a period of 60 days immediately prior to shipment, and no evidence of BTV transmission has been detected;

AND

6. were protected from attack from Culicoides likely to be competent BTV vectors during transportation to the place of shipment; or

7. were vaccinated 60 days before shipment or had antibodies against all serotypes whose presence in the zones of transit has been demonstrated through a surveillance and monitoring programme in accordance with Appendix 3.8.1.

Article 2.2.13.9.

When importing from BTV free countries or zones, Veterinary Administrations should require:

the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   a) were kept in a BTV free country or zone for at least 60 days before commencement of, and during, collection of the semen; or
Appendix XI (contd)

b) were subjected to a serological test according to the Terrestrial Manual to detect antibody to the BTV group, between 21 and 60 days after the last collection for this consignment, with negative results; or

c) were subjected to an agent identification test according to the Terrestrial Manual on blood samples collected at commencement and conclusion of, and at least every 7 days (virus isolation test) or at least every 28 days (PCR test) during, semen collection for this consignment, with negative results;

2. the semen was collected, processed and stored in conformity with Appendix 3.2.1.

Article 2.2.13.10.

When importing from BTV seasonally free zones, Veterinary Administrations should require:

for semen of ruminants and other BTV susceptible herbivores

the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   
a) were kept during the BTV seasonally free period in a seasonally free zone for at least 60 days before commencement of, and during, collection of the semen; or

   b) were subjected to a serological test according to the Terrestrial Manual to detect antibody to the BTV group, with negative results, at least every 60 days throughout the collection period and between 21 and 60 days after the final collection for this consignment; or

   c) were subjected to an agent identification test according to the Terrestrial Manual on blood samples collected at commencement and conclusion of, and at least every 7 days (virus isolation test) or at least every 28 days (PCR test) during, semen collection for this consignment, with negative results;

2. the semen was collected, processed and stored in conformity with Appendix 3.2.1.

Article 2.2.13.11.

When importing from BTV infected countries or zones, Veterinary Administrations should require:

for semen of ruminants and other BTV susceptible herbivores

the presentation of an international veterinary certificate attesting that:

1. the donor animals:
   
   a) were protected from attack from Culicoides likely to be competent BTV vectors for at least 60 days before commencement of, and during, collection of the semen; or

   b) were subjected to a serological test according to the Terrestrial Manual to detect antibody to the BTV group, with negative results, at least every 60 days throughout the collection period and between 21 and 60 days after the final collection for this consignment; or
Appendix XI (contd)

c) were subjected to an agent identification test according to the *Terrestrial Manual* on blood samples collected at commencement and conclusion of, and at least every 7 days (virus isolation test) or at least every 28 days (PCR test) during, semen collection for this consignment, with negative results;

2. the semen was collected, processed and stored in conformity with Appendix 3.2.1.

Article 2.2.13.12.

Regardless of the bluetongue status of the *exporting country*, *Veterinary Administrations* of *importing countries* should require:

for *in vivo* derived bovine embryos/oocytes

the presentation of an *international veterinary certificate* attesting that the embryos/oocytes were collected, processed and stored in conformity with the provisions of Appendix 3.3.1. or Appendix 3.3.3., as relevant.

Article 2.2.13.13.

When importing from BTV free countries or zones, *Veterinary Administrations* should require:

for *in vivo* derived embryos of ruminants (other than bovines) and other BTV susceptible herbivores

the presentation of an *international veterinary certificate* attesting that:

1. the donor females:
   a) were kept in a BTV free country or zone for at least the 60 days prior to, and at the time of, collection of the embryos; or
   b) were subjected to a serological test according to the *Terrestrial Manual* to detect antibody to the BTV group, between 21 and 60 days after collection, with negative results; or
   c) were subjected to an agent identification test according to the *Terrestrial Manual* on a blood sample taken on the day of collection, with negative results;

2. the embryos were collected, processed and stored in conformity with Appendix 3.3.1.

Article 2.2.13.14.

When importing from BTV seasonally free zones, *Veterinary Administrations* should require:

for *in vivo* derived embryos/oocytes of ruminants (other than bovines) and other BTV susceptible herbivores and for *in vitro* produced bovine embryos

the presentation of an *international veterinary certificate* attesting that:

1. the donor females:
   a) were kept during the seasonally free period in a seasonally free zone for at least 60 days before commencement of, and during, collection of the embryos/oocytes; or
Appendix XI (contd)

b) were subjected to a serological test according to the *Terrestrial Manual* to detect antibody to the BTV group, between 21 and 60 days after collection, with negative results; or

c) were subjected to an agent identification test according to the *Terrestrial Manual* on a blood sample taken on the day of collection, with negative results;

2. the embryos/oocytes were collected, processed and stored in conformity with Appendix 3.3.1.

**Article 2.2.13.15.**

When importing from BTV infected countries or zones, Veterinary Administrations should require:

for *in vivo* derived embryos/oocytes of ruminants (other than bovines) and other BTV susceptible herbivores and for *in vitro* produced bovine embryos

the presentation of an *international veterinary certificate* attesting that:

1. the donor females:
   
a) were protected from attack from *Culicoides* likely to be competent BTV vectors for at least 60 days before commencement of, and during, collection of the embryos/oocytes; or

b) were subjected to a serological test according to the *Terrestrial Manual* to detect antibody to the BTV group, between 21 and 60 days after collection, with negative results; or

   
c) were subjected to an agent identification test according to the *Terrestrial Manual* on a blood sample taken on the day of collection, with negative results;

2. the embryos/oocytes were collected, processed and stored in conformity with Appendix 3.3.1.

**Article 2.2.13.16.**

**Protecting animals from *Culicoides* attack**

When transporting animals through BTV infected countries or zones, Veterinary Administrations should require strategies to protect animals from attack from *Culicoides* likely to be competent BTV vectors during transport, taking into account the local ecology of the vector.

Potential risk management strategies include:

1. treating animals with chemical repellents prior to and during transportation;

2. loading, transporting and unloading animals at times of low vector activity (i.e. bright sunshine, 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 shadecloth;
5. monitoring for vectors at common stopping and offloading points to gain information on seasonal variations;

6. using historical, ongoing and/or BTV modelling information to identify low risk ports and transport routes.

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APPENDIX 3.X.X.

GUIDELINES FOR THE SURVEILLANCE OF BLUETONGUE

Article 3.X.X.1.

Introduction

This Appendix defines the principles and provides a guide for the surveillance of bluetongue (BT) in accordance with Appendix 3.8.1., applicable to countries seeking recognition for a declared BT status, with or without the use of vaccination. This may be for the entire country, zone or compartment. Guidance for countries seeking free status following an outbreak and for the maintenance of BT status is also provided. This Appendix complements Chapter 2.2.13.

BT is a vector-borne infection transmitted by different species of Culicoides insects in a range of ecosystems. An important component of BT epidemiology is vectorial capacity which provides a measure of disease risk that incorporates vector competence, abundance, biting rates, survival rates and extrinsic incubation period. However, methods and tools for measuring some of these vector factors remain to be developed, particularly in a field context. Therefore, surveillance for BT should focus on transmission in domestic ruminants.

Susceptible wild ruminant populations should be included in surveillance only if necessary for trade.

The impact and epidemiology of BT differ widely in different regions of the world and therefore it is impossible to provide specific guidelines for all situations. It is incumbent upon Member Countries to provide scientific data that explain the epidemiology of BT in the region concerned and adapt the surveillance strategies for defining their infection status (free, endemic or area of potential spread) to the local conditions. There is considerable latitude available to Member Countries to justify their infection status at an acceptable level of confidence.

Surveillance for BT should be in the form of a continuing programme.

Article 3.X.X.2.

Case definition

For the purposes of surveillance, a case refers to an animal infected with BT virus (BTV).

For the purposes of international trade, a difference must be made between a case as defined below and an animal that is potentially infectious to vectors. The conditions for trade are defined in Chapter 2.2.13 of the Terrestrial Code.

The purpose of surveillance is the detection of virus circulation in a country or zone and not the status of an individual animal or herds. Surveillance deals not only with the occurrence of clinical signs caused by BTV, but also with the presence of infection with BTV in the absence of clinical signs.

The following defines the occurrence of BTV infection:

1. BTV has been isolated and identified as such from an animal or a product derived from that animal,
Appendix XII (contd)

2. viral antigen or viral RNA specific to one or more of the serotypes of BTV has been identified in samples from one or more animals showing clinical signs consistent with BT, or epidemiologically linked to a confirmed or suspected case, or giving cause for suspicion of previous association or contact with BTV, or

3. antibodies to structural or nonstructural proteins of BTV that are not a consequence of vaccination have been identified in one or more animals showing clinical signs consistent with BT, or epidemiologically linked to a confirmed or suspected case, or giving cause for suspicion of previous association or contact with BTV.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 3.X.X.3.

General conditions and methods

1. A surveillance system in accordance with Appendix 3.8.1. should be under the responsibility of the Veterinary Administration. In particular:

   a) a formal and ongoing system for detecting and investigating outbreaks of disease should be in place;

   b) a procedure should be in place for the rapid collection and transport of samples from suspect cases of BT to a laboratory for BT 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 BT surveillance programme should:

   a) include an early warning system for reporting suspicious cases. Farmers and workers, who have day-to-day contact with domestic ruminants, as well as diagnosticians, should report promptly any suspicion of BT 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 Administration. 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 BTV. 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 BT should be investigated immediately and samples should be taken and submitted to an approved 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.

With regards to BT, compartment refers to establishments where animals are kept in a confirmed vector free environment to prevent BTV infection. Generally, the conditions to prevent exposure of susceptible animals to BTV infected vectors will be difficult to apply. However, under specific situations like artificial insemination centres or quarantine stations such conditions may be met. The testing requirements for animals kept in these facilities are described in Articles 2.2.13.11 and 2.2.13.15.
Appendix XII (contd)

Article 3.X.X.4.

Surveillance strategies

The target population for surveillance aimed at identification of disease and/or infection should cover susceptible domestic ruminants within the country, zone or compartment. Active and passive surveillance for BTV 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.

The strategy employed may be based on randomised sampling requiring surveillance consistent with demonstrating the absence of BTV 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 may be followed up with virological methods as appropriate.

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 may be used concurrently to define the BTV status of targeted populations.

A country should justify the surveillance strategy chosen as being adequate to detect the presence of BTV infection in accordance with Appendix 3.8.1. 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. sheep). Similarly, virological and serological testing may be targeted to species that rarely show clinical signs (e.g. cattle).

In vaccinated populations, serological and virological surveillance is necessary to detect the BTV types circulating to ensure that all circulating types are included in the vaccination programme.

If a Member Country wishes to declare freedom from BTV 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 and expected prevalence determine the level of confidence in the results of the survey. The applicant country must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Appendix 3.8.1. 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.
Appendix XII (contd)

The principles involved in surveillance for disease/infection are technically well defined. The design of surveillance programmes to prove the absence of BTV 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.

1. Clinical surveillance

Clinical surveillance aims at the detection of clinical signs of BT at the flock/herd level. Whereas significant emphasis is placed on the diagnostic value of mass serological screening, surveillance based on clinical inspection should not be underrated, particularly during a newly introduced infection. In sheep and occasionally goats, clinical signs may include oedema, hyperaemia of mucosal membranes, coronitis and cyanotic tongue.

BT suspects detected by clinical surveillance should always be confirmed by laboratory testing.

2. Serological surveillance

An active programme of surveillance of host populations to detect evidence of BTV transmission is essential to establish BTV status in a country or zone. Serological testing of ruminants is one of the most effective methods of detecting the presence of BTV. The species tested depends on the epidemiology of BTV infection, and the species available, in the local area. Cattle are usually the most sensitive indicator species.

Surveillance may include serological surveys, for example abattoir surveys, the use of sentinel animals, or a combination of methods.

The objective of serological surveillance is to detect antibodies against BTV using tests prescribed in the *Terrestrial Manual*. Positive BTV antibody tests results can have four possible causes:

- a) natural infection with BTV,
- b) vaccination against BTV,
- 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 survey purposes for BTV surveillance. However, the principles of survey design described in these guidelines and the requirements for a statistically valid survey for the presence of BTV infection should not be compromised.

The results of random or targeted serological surveys are important in providing reliable evidence that no BTV infection is present in a country, zone or compartment. It is, therefore, essential that the survey is thoroughly documented.

Serological surveillance in a free zone should target those areas that are at highest risk of BTV 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 BTV infection, either random or targeted sampling is suitable to select herds and/or animals for testing.
Appendix XII (contd)

A surveillance zone within a free country or zone should separate it from a potentially infected country or zone. Serological surveillance in a free country or zone should be carried out over an appropriate distance from the border with a potentially infected country or zone, based upon geography, climate, history of infection and other relevant factors.

Serological surveillance in infected zones will identify changes in the boundary of the zone, and can also be used to identify the BTV types circulating. In view of the epidemiology of BTV infection, either random or targeted sampling is suitable.

3. Virological surveillance

Isolation and genetic analysis of samples of BTV 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 herds

Sentinel herds are a form of targeted surveillance with a prospective study design. They are the preferred strategy for BTV surveillance. They comprise groups of unexposed animals managed at fixed locations and sampled regularly to detect new BTV infections.

The primary purpose of a sentinel herd programme is to detect BTV infections occurring at a particular place, for instance sentinel groups may be located on the usual boundaries of infected zones to detect changes in distribution of BTV. In addition, sentinel herd programmes allow incidence rates to be determined and the timing of infections to be observed.

A sentinel herd programme should use animals of known source and history of exposure, control management variables such as use of insecticides 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 BTV 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 BTV infection. Cattle are the most appropriate sentinels but other domestic ruminant species may be used. The only feature distinguishing groups of sentinels should be their geographical location.

Sera from sentinel herd programmes should be stored methodically in a serum bank to allow retrospective studies to be conducted in the event of new serotypes being isolated.
Appendix XII (contd)

The frequency of sampling will depend on the reason for choosing the sampling site. In endemic areas, virus isolation will allow monitoring of the serotypes and genotypes of BTV 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. Sentinels in declared free zones add to confidence that BTV infections are not occurring unobserved. In such cases, sampling prior to and after the possible period of transmission is sufficient.

The definitive measure of a country or zone's BTV infection status is detection and identification of the viruses. If virus isolation is required, sentinels should be sampled at sufficiently frequent intervals to ensure that samples are collected during the period of viraemia.

5. Vector surveillance

BTV is transmitted between ruminant hosts by vector 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 species present in an area, their seasonal incidence and profile, and their 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 domestic ruminants, or the use of drop traps over ruminant animals.

The number of traps to be used in a vector surveillance system and the frequency of their use will depend on the availability of resources but is also dependent upon the size or ecological characteristics of the area to be surveyed.

The operation of vector surveillance sites at the same locations as sentinel herds 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 (e.g. the use of sentinel herds of domestic ruminants) are preferred to detect virus circulation.

Article 3.X.X.5.

Documentation of BTV infection free status

1. Countries declaring freedom from BTV infection for the country, zone or compartment

In addition to the general conditions described in Chapter 2.2.13. of the *Terrestrial Code*, a Member Country declaring freedom from BTV infection 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 Appendix, to demonstrate absence of BTV infection during the preceding 24 months in susceptible domestic ruminant populations. This requires the support of a laboratory able to undertake identification of BTV infection through virus detection and antibody tests described in the *Terrestrial Manual*. This surveillance should be targeted to non-vaccinated animals. Clinical surveillance may be effective in sheep while serological surveillance is more appropriate in cattle.

OIE Terrestrial Animal Health Standards Commission/September 2005
2. **Additional requirements for countries, zones or compartments that practise vaccination**

Vaccination to prevent the transmission of BTV may be part of a disease control programme. The level of flock or herd immunity required to prevent transmission will depend on the flock or herd size, composition (e.g. species) and density of the susceptible population. It is therefore impossible to be prescriptive. The vaccine must also comply with the provisions stipulated for BTV vaccines in the *Terrestrial Manual*. Based on the epidemiology of BTV infection in the country, zone or compartment, it may be that a decision is reached to vaccinate only certain species or other subpopulations.

In countries or zones that practice vaccination there is a need to perform virological and serological tests to ensure the absence of virus circulation. These tests should be performed on non-vaccinated subpopulations or on sentinels. The tests have to be repeated at appropriate intervals according to the purpose of the surveillance programme. For example, longer intervals may be adequate to confirm endemicity, while shorter intervals may allow on-going demonstration of absence of transmission.

**Article 3.X.X.6.**

**The use and interpretation of serological and virus detection tests**

1. **Serological testing**

Ruminants infected with BTV produce antibodies to structural and non-structural viral proteins, as do animals vaccinated with current modified live virus vaccines. Antibodies to the BTV serogroup antigen are detected with high sensitivity and specificity by competitive ELISA (c-ELISA) and to a lesser extent by AGID as described in the *Terrestrial Manual*. Positive c-ELISA results can be confirmed by neutralization assay to identify the infecting serotype (s), however BTV infected ruminants can produce neutralizing antibodies to serotypes of BTV other than those to which they were exposed (false positive results), especially if they have been infected with multiple serotypes.

2. **Virus detection**

The presence of BTV in ruminant blood and tissues can be detected by virus isolation or polymerase chain reaction (PCR) as described in the *Terrestrial Manual*.

Interpretation of positive and negative results (both true and false) differs markedly between these tests because they detect different aspects of BTV infection, specifically (1) infectious BTV (virus isolation) and (2) nucleic acid (PCR). The following are especially relevant to interpretation of PCR assays:

a) The nested PCR assay detects BTV nucleic acid in ruminants long after the clearance of infectious virus. Thus positive PCR results do not necessarily coincide with active infection of ruminants. Furthermore, the nested PCR assay is especially prone to template contamination, thus there is considerable risk of false positive results.

b) PCR procedures other than real time PCR allow sequence analysis of viral amplicons from ruminant tissues, insect vectors or virus isolates. These sequence data are useful for creating data bases to facilitate important epidemiological studies, including the possible distinction of field and vaccine virus strains of BTV, genotype characterization of field strains of BTV, and potential genetic divergence of BTV relevant to vaccine and diagnostic testing strategies.

It is essential that BTV isolates are sent regularly to the OIE Reference Laboratories for genetic and antigenic characterization.
Appendix XII (contd)

**Figure 1**
Application of laboratory tests in serological surveillance

**Figure 2**
Application of laboratory tests in virological surveillance
CHAPTER 2.3.13.

BOVINE SPONGIFORM ENCEPHALOPATHY

Article 2.3.13.1.

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 Administrations 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) gelatin 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 were subject to passed ante-mortem and post-mortem inspections and were not suspect or confirmed BSE cases; and which has been prepared in a manner to avoid contamination with tissues listed in Article 2.3.13.13.;
   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 Administrations 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.

Article 2.3.13.2.

The BSE risk status of the cattle population of a country, zone or compartment should be determined on the basis of the following criteria:
Appendix XIII (contd)

1. the outcome of a risk assessment (which is reviewed annually), based on Section 1.3., identifying all potential factors for BSE occurrence and their historic perspective:

   a) Release assessment

   Release assessment consists of assessing the likelihood that the BSE agent has been introduced into the cattle population from a pre-existing agent in the indigenous ruminant population or via commodities potentially contaminated with the BSE agent, through a consideration of the following:

   i) the presence or absence of animal TSE agents in the country, zone or compartment and, if present, evidence regarding their prevalence based on the outcomes of surveillance;

   ii) meat-and-bone meal or greaves from the indigenous ruminant population;

   iii) imported meat-and-bone meal or greaves;

   iv) imported live ruminants;

   v) imported animal feed and feed ingredients;

   vi) imported products of ruminant origin for human consumption, which may have contained tissues listed in Article 2.3.13.13. and may have been fed to cattle;

   vii) imported products of ruminant origin for in vivo use in cattle.

   The results of any surveillance and other epidemiological investigation into the disposition of the commodities identified above (especially surveillance for BSE conducted on the cattle population) relevant to the 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 exposure of the BSE agent to cattle, through a consideration of the following:

   i) 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;

   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;
Appendix XIII (contd)

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 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 Appendix 3.8.4.;

3. the compulsory notification and investigation of all cattle showing clinical signs consistent with BSE;

4. the examination in an approved laboratory of brain or other tissues collected within the framework of the aforementioned surveillance and monitoring system.

When the risk assessment (which takes into account the surveillance referred to in the release and exposure assessments above) demonstrates negligible risk, the country should conduct Type B surveillance in accordance with Appendix 3.8.4.

When the risk assessment (which takes into account the surveillance referred to in the release and exposure assessments above) demonstrates non-negligible risk, fails to demonstrate negligible risk, the country should conduct Type A surveillance in accordance with Appendix 3.8.4.

Article 2.3.13.3.

Negligible BSE risk

Commodities from the cattle population of a country, zone or compartment pose a negligible risk of transmitting the BSE agent, should the following conditions be met:

1. a risk assessment, as described in point 1) of Article 2.3.13.2., has been conducted in order to identify the historical and existing risk factors, and the country has demonstrated that appropriate generic specific measures have been taken for the relevant period of time defined below to manage all risks each risk identified;

2. the country has demonstrated that Type B surveillance, in accordance with Appendix 3.8.4, 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 any 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 2.3.13.2. 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 meat-and-bone meal or greaves derived from ruminants has not been fed to ruminants;
Appendix XIII (contd)

OR

b) the last indigenous case of BSE was reported more than 7 years ago, any indigenous case of BSE was born more than 8 years ago; and

i) the criteria in points 2) to 4) of Article 2.3.13.2. 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 meat-and-bone meal and greaves derived from ruminants has not been fed to ruminants; and

iii) all BSE cases, as well as:

- 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 2.3.13.4.

Controlled BSE risk

Commodities from the cattle population of a country, zone or compartment pose a controlled risk of transmitting the BSE agent, should the following conditions be met:

1. a risk assessment, as described in point 1) of Article 2.3.13.2., has been conducted in order to identify the historical and existing risk factors, and the country has demonstrated that appropriate measures are being taken, but have not been taken for the relevant period of time to manage all risks identified.

the country has not demonstrated that appropriate generic measures have been taken for the relevant period of time defined below to manage all risks identified.

2. the country has demonstrated that Type A surveillance in accordance with Appendix 3.8.4. is in place; Type B surveillance may replace Type A surveillance once the relevant points target, in accordance with Table 1, is met.

3. EITHER

a) there has been no case of BSE or any case of BSE has been demonstrated to have been imported and has been completely destroyed, the criteria in points 2) to 4) of Article 2.3.13.2. are complied with, and it can be demonstrated, through an appropriate level of control and audit, that meat-and-bone meal and greaves derived from ruminants has not been fed to ruminants, but at least one of the following two conditions applies:
Appendix XIII (contd)

i) the criteria in points 2) to 4) of Article 2.3.13.2. 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;

OR

b) there has been an indigenous case of BSE reported, the criteria in points 2) to 4) of
Article 2.3.13.2. are complied with, and it can be demonstrated, through an appropriate level of
control and audit that meat-and-bone meal and greaves derived from ruminants have not been fed
to ruminants, but at least one of the following two conditions applies:

i) the criteria in points 2) to 4) of Article 2.3.13.2. have not been complied with for 7 years;

ii) it cannot be demonstrated that controls over the feeding of meat-and-bone meal and greaves
derived from ruminants to ruminants have been in place for 8 years;

AND

iii) all BSE cases, as well as:

- 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 2.3.13.5.

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 2.3.13.6.

When importing from a country, zone or compartment posing a negligible BSE risk, Veterinary
Administrations should require:

for all commodities from cattle not listed in point 1) of Article 2.3.13.1.

the presentation of an international veterinary certificate attesting that the country, zone or compartment
complies with the conditions in Article 2.3.13.3.
Appendix XIII (contd)

Article 2.3.13.7.

When importing from a country, zone or compartment posing a controlled BSE risk, Veterinary Administrations should require:

for cattle

the presentation of an international veterinary certificate attesting that:

1. the country, zone or compartment complies with the conditions in Article 2.3.13.4.;

2. cattle selected for export are identified by a permanent identification system enabling them to be traced back to the dam and herd of origin, and are not exposed cattle as described in point 3) b) iii) of Article 2.3.13.4.;

3. in the case of a country, zone or compartment with an indigenous case, 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 had been effectively enforced.

Article 2.3.13.8.

When importing from a country, zone or compartment with an undetermined BSE risk, Veterinary Administrations should require:

for cattle

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 the progeny of female cases, born within 2 years prior to or after clinical onset of the disease, and
   
   b) 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
   
   c) 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 enabling them to be traced back to the dam and herd of origin and are not the progeny of BSE suspect or confirmed females;
   
   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.
Appendix XIII (contd)

Article 2.3.13.9.

When importing from a country, zone or compartment posing a negligible BSE risk, Veterinary Administrations should require:

for fresh meat and meat products from cattle (other than those listed in point 1) of Article 2.3.13.1.)

the presentation of an international veterinary certificate attesting that:

1. the country, zone or compartment complies with the conditions in Article 2.3.13.3.;

2. the cattle from which the fresh meat and meat products were derived passed ante-mortem and post-mortem inspections; ante-mortem and post-mortem inspections were carried out on all cattle from which the fresh meat or meat products originate.

Article 2.3.13.10.

When importing from a country, zone or compartment posing a controlled BSE risk, Veterinary Administrations should require:

for fresh meat and meat products from cattle (other than those listed in point 1) of Article 2.3.13.1.)

the presentation of an international veterinary certificate attesting that:

1. the country, zone or compartment complies with the conditions in Article 2.3.13.4.;

2. the cattle from which the fresh meat and meat products were derived passed ante-mortem and post-mortem inspections; ante-mortem and post-mortem inspections were carried out on all cattle from which the fresh meat and meat products originate;

3. cattle from which the fresh meat and meat products destined for export originate 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 pitting process;

4. the fresh meat and meat products do not contain 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 Article 2.3.13.13.,

   b) mechanically separated meat from the skull and vertebral column from cattle over 30 months of age,

   all of which have been completely removed in a manner to avoid contamination of the fresh meat and meat products.

Article 2.3.13.11.

When importing from a country, zone or compartment with an undetermined BSE risk, Veterinary Administrations should require:
Appendix XIII (contd)

for fresh meat and meat products from cattle (other than those listed in point 1) of Article 2.3.13.1.)

the presentation of an international veterinary certificate attesting that:

1. the cattle from which the fresh meat and meat products originate:
   a) are not suspect or confirmed BSE cases;
   b) have not been fed meat-and-bone meal or greaves derived from ruminants;
   c) were subjected to passed ante-mortem and post-mortem inspections;
   d) 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 do not contain 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 Article 2.3.13.13.,
   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,

all of which have been completely removed in a manner to avoid contamination of the fresh meat and meat products.

Article 2.3.13.12.

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 2.3.13.4. and 2.3.13.5. should not be traded between countries.

Article 2.3.13.13.

1. From cattle of any age originating from a country, zone or compartment defined in Articles 2.3.13.4. and 2.3.13.5., 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, and protein products derived thereof. Food, feed, fertilisers, cosmetics, pharmaceuticals or medical devices prepared using these commodities 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 2.3.13.4., 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, vertebral column and derived protein products. Food, feed, fertilisers, cosmetics, pharmaceuticals or medical devices prepared using these commodities 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 2.3.13.5., 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, vertebral column and derived protein products. Food, feed, fertilisers, cosmetics, pharmaceuticals or medical devices prepared using these commodities should also not be traded.

Article 2.3.13.14.

Veterinary Administrations of importing countries should require:

for gelatin and collagen prepared from bones and intended for food or feed, cosmetics, pharmaceuticals including biologicals, or medical devices

the presentation of an international veterinary certificate attesting that the commodities came from:

1. a country, zone or compartment posing a negligible BSE risk; or
2. a country, zone or compartment posing a controlled BSE risk; and
   a) skulls and vertebrae (except tail vertebrae) have been excluded;
   b) the bones have been subjected to a process which includes all the following steps:
      i) pressure washing (degreasing),
      ii) acid demineralisation,
      iii) prolonged alkaline treatment,
      iv) filtration,
      v) sterilisation at ≥138°C for a minimum of 4 seconds,

   or to an equivalent process in terms of infectivity reduction.

Article 2.3.13.15.

Veterinary Administrations of importing countries should require:

for tallow and dicalcium phosphate (other than protein-free tallow as defined in Article 2.3.13.1.) intended for food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices

the presentation of an international veterinary certificate attesting that it originates from:

1. a country, zone or compartment posing a negligible BSE risk; or
2. a country, zone or compartment posing a controlled BSE risk, and
   it originates from cattle which have been subjected to passed ante-mortem and post-mortem inspections, and has not been prepared using the tissues listed in points 1 and 2 of Article 2.3.13.13.
Appendix XIII (contd)

Article 2.3.13.16.

Veterinary Administrations of importing countries should require:

for tallow derivatives (other than those made from protein-free tallow as defined in Article 2.3.13.1.) intended for food, feed, fertilisers, cosmetics, pharmaceuticals including biologicals, or medical devices

the presentation of an international veterinary certificate attesting that:

1. they originate from a country, zone or compartment posing a negligible BSE risk; or
2. they have been produced by hydrolysis, saponification or transesterification using high temperature and pressure.

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APPENDIX 3.8.4.

SURVEILLANCE FOR BOVINE SPONGIFORM ENCEPHALOPATHY

Article 3.8.4.1.

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;
   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 2.3.13.2. 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:
Appendix XIV (contd)

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 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. All countries should sample at least three of the four subpopulations. This approach is consistent with Appendix 3.8.1. on general guidelines for animal health surveillance.

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 3.8.4.2.

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 countries 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, particularly cattle over 30 months of age, is the one exhibiting the highest prevalence. The recognition greatly depends on the owner’s awareness and observation of suspect animals. The reporting of these suspect animals when at the farm will depend on the owner’s motivation based on cost and socio-economic repercussions. 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 2.3.13.2), 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 countries 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 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 countries 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 countries 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).

Within each of the above subpopulations, countries may wish to target cattle identifiable as imported from countries or zones not free from BSE, cattle which have consumed potentially contaminated feedstuffs from countries or zones not free from BSE, offspring of BSE affected cows and cattle which have consumed feedstuffs potentially contaminated with other TSE agents.

When establishing a surveillance strategy, authorities must take into account inherent difficulties of obtaining samples on farm. These difficulties include higher cost, necessity for education and motivation of owners, counteracting potentially negative socio-economic implication. Authorities must find ways to overcome these difficulties.

Article 3.8.4.3.

4) Implementation of Type A surveillance

In order to implement efficiently a surveillance strategy for BSE, a country must use good quality data (or reliable estimates) documented records or reliable estimates of concerning the age distribution of its adult cattle population and the number of cattle tested for BSE stratified by age and by subpopulation within the country, zone or compartment. The application of the following procedure will allow the detection of BSE at a prevalence of at least one case per 100,000 in the adult cattle population, at a confidence level of 95% in the country, zone or compartment of concern.

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.
Appendix XIV (contd)

A country should design its surveillance strategy 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 appendix were obtained by applying the following factors to a statistical model:

a) the design prevalence for Type A or Type B surveillance of one case per 100,000 of the adult cattle population;

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 clinical 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:

a) cattle population numbers stratified by age;

b) the number of cattle tested for BSE stratified by age and by subpopulation.

This Appendix 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 country 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. **Maintenance (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 2.3.13.3) 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 2.3.13.4), following the achievement of the relevant points target using Type A surveillance, to maintain confidence in the knowledge gained through Type A surveillance.

   For countries which have demonstrated through risk assessment (including surveillance) that they meet the requirements for ‘negligible risk’, should continue at a reduced maintenance level.

   In order to implement efficiently a maintenance surveillance strategy for BSE, a country must use good quality data (or reliable estimates) concerning the age distribution of its adult cattle population and the number of cattle tested for BSE stratified by age and by subpopulation. The application of the following procedure will allow the detection of BSE prevalence of at least one case per 50,000 in the adult cattle population, at a confidence level of 95% in the country, zone or compartment of concern. This Appendix utilises Tables 1 and 2 to determine a desired surveillance point target and the point values of surveillance samples collected.

   Maintenance surveillance should focus on the higher prevalence subpopulations (especially clinical suspects). The number of clinical suspect samples taken annually should approximate the number of samples taken annually from clinical suspect cases during the time taken to reach the country, zone or compartment’s BSE status (to a maximum of 7 years).

   Article 3.8.4.4.

1. **Selecting the points target**

   The desired surveillance points target should be selected from Table 1, which shows target points for adult cattle populations of different sizes. A country’s The size of the adult cattle population size 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. The target depends on the design prevalence chosen by the country.

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47 DP (design prevalence) is used to determine the size of a testing survey expressed in terms of target points. If the actual prevalence is greater than the selected design prevalence, the survey is highly likely to detect disease.
Table 1  Points targets for different adult cattle population sizes in a country, zone or compartment which has not identified any BSE cases

<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>≥ 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>
</tbody>
</table>

DP is the maximum possible prevalence or “design prevalence”.

2. 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 Appendix 3.8.1. 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.

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’.

In addition, Countries should sample at least three of the four subpopulations.

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.
### 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(^1)</th>
<th>Fallen stock(^2)</th>
<th>Casualty slaughter(^3)</th>
<th>Clinical suspect(^4)</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>

\(^1\) See point 4) of Article 3.8.4.2.
\(^2\) See point 3) of Article 3.8.4.2.
\(^3\) See point 2) of Article 3.8.4.2.
\(^4\) See point 1) of Article 3.8.4.2.

Surveillance points remain valid for 7 years (the 95\(^{th}\) percentile of the incubation period).

**Article 3.8.4.5.**

To monitor the evolution of BSE in a country, zone or compartment once it is detected

To monitor the evolution of BSE in a country, zone or compartment once it is detected, a more intensive sampling method needs to be used to determine disease prevalence. For countries that have determined that BSE exists within their cattle population, the goal of surveillance shifts from one of detection to one of monitoring the extent and evolution of the disease, and monitoring the effectiveness of control measures such as feed bans and policies for the removal of specified risk materials.
CHAPTER 2.6.7.
CLASSICAL SWINE FEVER

Article 2.6.7.1.

The pig is the only natural host for classical swine fever (CSF) virus. The definition of pigs includes all varieties of *Sus scrofa*, both domestic breeds and wild boar. A distinction is made between farmed and permanently captive pigs, and free-living pigs. Farmed and permanently captive pigs of any breed will hereafter be referred to as domestic pigs. Free-living pigs of any breed will hereafter be referred to as wild pigs. Extensively kept pigs may fall into either of these categories or may alternate between the two. 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).

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 7-10 days, and are usually infective between post-infection days 5 and 14, but up to 3 months in cases of chronic infections.

Standards for diagnostic tests and vaccines are described in the *Terrestrial Manual*.

Article 2.6.7.2.

The CSF status of a country or zone can only be determined after considering the following criteria both 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 country, and all clinical signs suggestive of CSF should be subjected to 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 Administration should have current knowledge of, and authority over, all domestic establishments containing pigs in the whole country, zone or compartment;
5. the Veterinary Administration should have current knowledge about the population and habitat of wild pigs in the whole country or zone.

Article 2.6.7.3.

For the purposes of the *Terrestrial Code*:

‘CSF infected establishment’ means a domestic pig holding in which the presence of the infection has been confirmed by field and/or laboratory investigations.
Appendix XV (contd)

'Country, zone or compartment with CSF infection in domestic pigs' means a country, zone or compartment containing a CSF-infected establishment.

The size and limits of a CSF domestic pig control area must be based on the control measures used and the presence of natural and administrative boundaries, as well as an assessment of the risks for disease spread.

Article 2.6.7.4.

Country or zone Country, zone or compartment free of CSF in domestic and wild pigs

1. Historically free status

A country or zone country, zone or compartment may be considered free from the disease in domestic and wild pigs after conducting a risk assessment as referred to in Article 2.6.7.2, but without formally applying a specific surveillance programme (historical freedom) if the country or zone complies with the provisions of Appendix 3.8.48.

2. Free status as a result of an eradication a specific surveillance programme

A country or zone country, zone or compartment which does not meet the conditions of point 1) above may be considered free from CSF in domestic and wild pigs after the conducting of a risk assessment as referred to in Article 2.6.7.2, and surveillance in accordance with Appendix 3.8.8, is in place, and when:

a) CSF is a notifiable disease;

AND EITHER

b) no outbreak has been observed in domestic pigs for at least 12 months; or

b) bis where a stamping-out policy without vaccination has been practised for CSF control, no outbreak has been observed in domestic pigs for at least 6 months; or

c) where a stamping-out policy with vaccination is practised, either

i) 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;

c) bis where a vaccination strategy is practised has been adopted, with or without a stamping-out policy,

i) vaccination against CSF has been banned in all domestic pigs in the country or zone country, zone or compartment for at least 12 months one year, unless there are validated means of distinguishing between vaccinated and infected pigs;


Appendix XV (contd)

ii) if vaccination has been practised within the past 5 years, surveillance in accordance with Appendix 3.8.8. 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

d) based on surveillance in accordance with Appendix 3.8.8, CSF infection is not known to occur in the any wild pig population in the country, zone or compartment and surveillance of wild pigs indicates that there is no residual infection.

Article 2.6.7.5.

Country or zone free of CSF in domestic pigs but with a infection in the wild pig population

Requirements in point 2) a) to c)bis of Article 2.6.7.4. as relevant, are complied with. As but CSF infection is known to occur may be present in the wild pigs population, the following additional conditions are complied with for the free status are that in the country or zone:

1. a programme for the management of CSF in wild pigs is in place, and CSF wild pig control areas are delineated around every CSF case reported in wild pigs, 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. biosecurity measures are zoning or compartmentalisation is applied to prevent transmission of CSF from wild pigs to domestic pigs;

3. surveillance in accordance with Appendix 3.8.8. is carried out in the domestic pig population, with negative results.

Article 2.6.7.6.

Recovery of free status

Should a CSF outbreak occur in an establishment of a free country or zone country, zone or compartment (free in domestic and wild pigs, or free in domestic pigs only), the status of the country, or zone or compartment may be restored at least not less than 30 days after completion of a stamping-out policy where surveillance in accordance with Appendix 3.8.8. has been carried out with negative results, which should include the following measures:

1. a CSF domestic pig control area (including an inner protection area of at least 3 kilometre radius and an outer surveillance area of at least 10 kilometre radius) should be delineated around the outbreak, taking into account the control measures applied, the presence of natural and administrative boundaries, and an assessment of the risk of disease spread;
Appendix XV (contd)

2. all the pigs have been killed and their carcasses destroyed, and disinfection has been applied within the establishment;

3. in the protection area around a CSF outbreak:
   a) a risk assessment should be carried out to determine the likelihood of CSF infection in neighbouring establishments; when a significant risk is indicated, a stamping-out policy of all domestic pigs within a radius of at least 0.5 kilometre may be applied;
   b) an immediate clinical examination of all pigs in all pig establishments situated within the protection area has been carried out;

4. in the surveillance area around a CSF outbreak, all sick pigs should be subjected to laboratory tests for CSF;

5. surveillance in accordance with Appendix 3.8.8. has been carried out in all pig establishments that have been directly or indirectly in contact with the infected establishment and in all pig establishments located within the CSF domestic pig control area, demonstrating that these establishments are not infected;

6. measures aimed at preventing any virus spread by live pigs, pig semen and pig embryos, contaminated material, vehicles, etc. have been implemented.

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.

Article 2.6.7.7.

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 Appendix 3.8.8. 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, 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 3.6.4.1.;
5. imported wild pigs comply with the relevant requirements set forth in the present chapter.

A zoning compartmentalisation approach within the country or zone can only be adopted if there is a wild pig sub-population that is isolated through a biosecurity management system from other wild pigs.

**Article 2.6.7.8.**

When importing from countries or zones countries, zones or compartments free of CSF in domestic and wild pigs, Veterinary Administrations should require:

for domestic pigs

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 or zone country, zone or compartment free of CSF in domestic and wild pigs 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 validated means of distinguishing between vaccinated and infected pigs.

**Article 2.6.7.9.**

When importing from countries free of CSF in domestic pigs but with a wild pig population countries or zones free of CSF in domestic pigs but with infection in the wild pig population, Veterinary Administrations should require:

for domestic pigs

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 an establishment a free zone or compartment which is not located in a CSF wild pig control area as defined in Article 2.6.7.5., and has undergone surveillance to verify absence of CSF in accordance with Appendix 3.8.8.;
4. have had no contact with pigs introduced into the establishment during the past 40 days;
5. showed no clinical sign of CSF on the day of shipment.

**Article 2.6.7.10.**

When importing from countries or zones with CSF infection in domestic pigs, Veterinary Administrations should require:
Appendix XV (contd)

for domestic pigs

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;

2. were kept since birth or for the past 3 months, in an establishment a free compartment not situated in a CSF domestic or wild pig control area as defined in Article 2.6.7.5 and in Article 2.6.7.6;

3. were isolated in a quarantine station for at least 40 days;

4. were subjected during that period of quarantine to a virological test, and a serological test performed at least 21 days after entry into the quarantine station, with negative results;

5. showed no clinical sign of CSF on the day of shipment.

Article 2.6.7.11.

When importing from countries or zones free of CSF in domestic and wild pigs, Veterinary Administrations should require:

for wild pigs

the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of CSF on the day of shipment;

2. have been captured in a country or zone free from CSF in domestic and wild pigs;

3. have not been vaccinated against CSF, unless there are validated means 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 2.6.7.12.

When importing from countries or zones, zones or compartments free of CSF in domestic and wild pigs, Veterinary Administrations should require:

for semen of domestic pigs

the presentation of an international veterinary certificate attesting that:

1. the donor animals:

   a) were kept in a country or zone country, zone or compartment free of CSF in domestic and wild pigs since birth or for at least the past 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 Appendix 3.2.2.

Article 2.6.7.13.

When importing from countries or zones free of CSF in domestic pigs but with infection in the wild pig population, Veterinary Administrations should require:

for semen of domestic pigs

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 have been kept in an artificial insemination centre which is not located in a CSF wild pig control area and is regularly monitored to verify absence of CSF in accordance with Appendix 3.8.8;
   b) were isolated in the artificial insemination centre for at least 40 days prior to collection;
   c) 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 Appendix 3.2.2.

Article 2.6.7.14.

When importing from countries or zones considered infected with CSF in domestic pigs, Veterinary Administrations should require:

for semen of domestic pigs

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;
   a)bis showed no clinical sign of CSF on the day of collection of the semen and for the following 40 days;
   b) have not been vaccinated against CSF, and were subjected to a serological test performed at least 21 days after collection, with negative results;

2. the semen was collected, processed and stored in conformity with the provisions of Appendix 3.2.2.

Article 2.6.7.15.

When importing from countries, or zones or compartments free of CSF in domestic and wild pigs, Veterinary Administrations should require:
Appendix XV (contd)

for in vivo derived embryos of pigs

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 Appendix 3.3.1.

Article 2.6.7.16.

When importing from countries or zones free of CSF in domestic pigs but with infection in the wild pig population, Veterinary Administrations should require:

for in vivo derived embryos of pigs

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 were kept for at least 40 days prior to collection in an establishment which is not located in a CSF domestic or wild pig control area and is regularly monitored to verify absence of CSF in accordance with Appendix 3.8.8.;
   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 Appendix 3.3.1.

Article 2.6.7.17.

When importing from countries or zones considered infected with CSF in domestic pigs, Veterinary Administrations should require:

for in vivo derived embryos of pigs

the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) were kept in a compartment free of CSF in domestic pigs since birth or for at least 3 months prior to collection were kept for at least 40 days prior to collection in an establishment which is not located in a CSF domestic or wild pig control area and is regularly monitored to verify absence of CSF in accordance with Appendix 3.8.8.;
   b) showed no clinical sign of CSF on the day of collection of the embryos and for the following 24 40 days;
   c) have not been vaccinated against CSF and were subjected, with negative results, to a serological test performed at least 21 days after collection;
2. the embryos were collected, processed and stored in conformity with the provisions of Appendix 3.3.1.
Appendix XV (contd)

Article 2.6.7.18.

When importing from countries, or zones or compartments free of CSF in domestic and wild pigs, Veterinary Administrations should require:

for fresh meat of domestic pigs

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, or zone or compartment free of CSF in domestic and wild 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 and have been found free of any sign suggestive of CSF.

Article 2.6.7.19.

When importing from countries or zones free of CSF in domestic pigs but with infection in the wild pig population, Veterinary Administrations should require:

for fresh meat of domestic pigs

the presentation of an international veterinary certificate attesting that the entire consignment of meat comes from animals which:

1. were kept in a country, or zone or compartment free of CSF in domestic pigs since birth or for at least the past 3 months;
2. were kept in an establishment which was not located in a CSF wild pig control area and had undergone surveillance to verify absence of CSF in accordance with Appendix 3.8.8;
3. have been slaughtered in an approved abattoir not located in a CSF control area, have been subjected to ante-mortem and post-mortem inspections and have been found free of any sign suggestive of CSF.

Article 2.6.7.20.

When importing from countries or zones free of CSF in domestic and wild pigs, Veterinary Administrations should require:

for fresh meat of wild pigs

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 country or zone free of CSF in domestic and wild pigs;
   b) have been subjected to post-mortem inspection in an approved examination centre, and have been found free of any sign suggestive of CSF;
Appendix XV (contd)

and, if the zone where the animal has been killed is adjacent to a zone with infection in wild pigs:

2. a sample has been collected from every animal shot, and has been subjected to a virological test and a serological test for CSF, with negative results.

Article 2.6.7.21.

Veterinary Administrations of importing countries should require:

for 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

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 2.6.7.18., 2.6.7.19. or 2.6.7.20., as relevant;
   b) in a processing establishment:
      i) approved by the Veterinary Administration for export purposes;
      ii) regularly inspected by the Veterinary Authority;
      iii) not situated in a CSF control area;
      iv) processing only meat meeting the conditions laid down in Articles 2.6.7.18., 2.6.7.19. or 2.6.7.20., as relevant;

OR

2. have been processed in an establishment approved by the Veterinary Administration for export purposes and regularly inspected by the Veterinary Authority so as to ensure the destruction of the CSF virus in conformity with one of the procedures referred to in Article 3.6.4.2.

Article 2.6.7.22.

Veterinary Administrations of importing countries should require:

for products of animal origin (from pigs, but not derived from fresh meat) intended for use in animal feeding and for agricultural or industrial use

the presentation of an international veterinary certificate attesting that the products:

1. have been prepared:
   a) exclusively from products meeting the conditions laid down for fresh meat in Articles 2.6.7.18., 2.6.7.19. or 2.6.7.20., as relevant;
Appendix XV (contd)

b) in a processing establishment:
   i) approved by the *Veterinary Administration* for export purposes;
   ii) regularly inspected by the *Veterinary Authority*;
   iii) not situated in a CSF control area;
   iv) processing only products meeting the conditions laid down in point a) above;

OR

2. have been processed in an establishment approved by the *Veterinary Administration* for export purposes and regularly inspected by the *Veterinary Authority* so as to ensure the destruction of the CSF virus in conformity with one of the procedures referred to in Article 3.6.4.2.

   Article 2.6.7.23.

*Veterinary Administrations of importing countries* should require:

for bristles (from pigs)

the presentation of an *international veterinary certificate* attesting that the products:

1. come from a country, or *zone or compartment* free of CSF in domestic and wild pigs; or

2. have been processed in an establishment approved by the *Veterinary Administration* for export purposes and regularly inspected by the *Veterinary Authority* so as to ensure the destruction of the CSF virus.

   Article 2.6.7.24.

*Veterinary Administrations of importing countries* should require:

for litter and manure (from pigs)

the presentation of an *international veterinary certificate* attesting that the products:

1. come from a country, or *zone or compartment* free of CSF in domestic and wild pigs; or

2. come from establishments situated in a country or *zone free of CSF in domestic pigs but with infection in wild pigs, but not located in a CSF control area*; or

3. have been processed in an establishment approved by the *Veterinary Administration* for export purposes and regularly inspected by the *Veterinary Authority* so as to ensure the destruction of the CSF virus.
CHAPTER 2.7.12.

AVIAN INFLUENZA

Article 2.7.12.1.

1. For the purposes of this Terrestrial Code, 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 birds reared or kept in captivity 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’.

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. 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; or

   c) antibodies to H5 or H7 subtype of NAI virus that are not a consequence of vaccination have been detected in poultry. In the case of isolated serological positive results, NAI infection may be ruled out on the basis of a thorough epidemiological investigation that does not demonstrate further evidence of NAI infection.
Appendix XVI (contd)

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 Appendix 3.8.9.

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 2.7.12.2.

The NAI 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 NAI occurrence and their historic perspective;

2. 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;

3. 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 an NAI surveillance programme in accordance with Appendix 3.8.9.

Article 2.7.12.3.

**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 has been present in the country, *zone or compartment* for the past 12 months, based on surveillance in accordance with Appendix 3.8.9. 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 a previously free country, *zone or compartment*, 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 Appendix 3.8.9. 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 *specified* conditions specified in Article 2.7.12.19 or 2.7.12.20 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 Appendix 3.8.9. has been carried out during that three-month period.
Appendix XVI (contd)

Article 2.7.12.4.

HPNAI free country, zone or compartment

A country, zone or compartment may be considered free from HPNAI when it has been shown that HPNAI infection has not been present in the country, zone or compartment for the past 12 months, although its LPNAI status may be unknown, when, based on surveillance in accordance with Appendix 3.8.9., 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 zones or compartments depending on historical or geographical factors, industry structure, population data, or proximity to recent outbreaks.

If infection has occurred in a previously free country, zone or compartment, 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 Appendix 3.8.9. has been carried out during that three-month period.

Article 2.7.12.5.

When importing from an NAI free country, zone or compartment, Veterinary Administrations should require:

for live poultry (other than day-old poultry)

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 an NAI free country, zone or compartment since they were hatched or for the past 21 days;
3. the required surveillance has been carried out on the establishment within the past 21 days;
4. if vaccinated, the poultry have been vaccinated in accordance with Appendix 3.8.9., and the relevant information is attached.

Information concerning the vaccination status of the poultry (including the dates of vaccination, and the vaccine used should be included in the veterinary certificate.

Article 2.7.12.6.

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Administrations should require:

for live birds other than poultry

the presentation of an international veterinary certificate attesting that the birds:

1. showed no clinical sign of infection with a virus which would be considered NAI in poultry on the day of shipment;
2. 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;
Appendix XVI (contd)

3. were subjected to a diagnostic test 7 to 14 days prior to shipment to demonstrate freedom from infection with a virus which would be considered NAI in poultry;

4. are transported in new containers.

Article 2.7.12.7.

When importing from an NAI free country, zone or compartment, Veterinary Administrations should require:

for day-old live poultry

the presentation of an international veterinary certificate attesting that the poultry:

1. the poultry were kept in an NAI free country, zone or compartment since they were hatched;

2. the poultry were derived from parent flocks which had been kept in an NAI free country, zone or compartment for at least 21 days prior to and at the time of the collection of the eggs;

3. if the poultry or the parent flocks were vaccinated, vaccination was carried out in accordance with Appendix 3.8.9., and the relevant information is attached.

Information concerning the vaccination status of the poultry and the parent flocks (including the dates of vaccination, and the vaccine used) should be included in the veterinary certificate.

Article 2.7.12.8.

When importing from an HPNAI free country, zone or compartment, Veterinary Administrations should require:

for day-old live poultry

the presentation of an international veterinary certificate attesting that the poultry:

1. the poultry were kept in an HPNAI free country, zone or compartment since they were hatched;

2. the poultry were derived from parent flocks which had been kept in an 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 containers.

4. if the poultry or the parent flocks were vaccinated, vaccination was carried out in accordance with Appendix 3.8.9., and the relevant information is attached.

Information concerning the vaccination status of the poultry and the parent flocks (including the dates of vaccination, and the vaccine used) should be included in the veterinary certificate.
Article 2.7.12.9.

When importing from an NAI free country, \textit{zone or compartment}, \textit{Veterinary Administrations} should require:

\textbf{for hatching eggs}

the presentation of an \textit{international veterinary certificate} attesting that the eggs:

1. the eggs came from an NAI free country, \textit{zone or compartment};

2. the eggs were derived from parent flocks which had been kept in an NAI free country, \textit{zone or compartment} for at least 21 days prior to and at the time of the collection of the eggs.

3. if the parent flocks were vaccinated, vaccination was carried out in accordance with Appendix 3.8.9, and the relevant information is attached.

Information concerning the vaccination status of the parent flocks (including the dates of vaccination, and the vaccine used) should be included in the veterinary certificate.

Article 2.7.12.10.

When importing from a HPNAI free country, \textit{zone or compartment}, \textit{Veterinary Administrations} should require:

\textbf{for hatching eggs}

the presentation of an \textit{international veterinary certificate} attesting that the eggs:

1. the eggs came from an HPNAI free country, \textit{zone or compartment};

2. the eggs were derived from parent flocks which had been kept in an 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 Article 3.4.1.7) and are transported in new packing material;

4. if the parent flocks were vaccinated, vaccination was carried out in accordance with Appendix 3.8.9, and the relevant information is attached.

Information concerning the vaccination status of the parent flocks (including the dates of vaccination, and the vaccine used) should be included in the veterinary certificate.

Article 2.7.12.11.

When importing from an NAI free country, \textit{zone or compartment}, \textit{Veterinary Administrations} should require:

\textbf{for eggs for human consumption}

the presentation of an \textit{international veterinary certificate} attesting that the eggs come from an NAI free country, \textit{zone or compartment}.
Appendix XVI (contd)

Article 2.7.12.12.

When importing from a HPNAI free country, zone or compartment, Veterinary Administrations should require:

for eggs for human consumption

the presentation of an international veterinary certificate attesting that the eggs:

1. come from a HPNAI free country, zone or compartment;

2. come from establishments in which there has been no evidence of NAI in the past 21 days;

3. have had their surfaces sanitised (in accordance with Article 3.4.1.7) and are transported in new packing material.

Article 2.7.12.13.

When importing from an NAI free country, zone or compartment, Veterinary Administrations should require:

for egg products

the presentation of an international veterinary certificate attesting that the egg products come from, and were processed in, an NAI free country, zone or compartment.

Article 2.7.12.14.

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Administrations should require:

for egg products

the presentation of an international veterinary certificate attesting that the egg products:

1. are derived from eggs which meet the requirements of Articles 2.7.12.9., 2.7.12.10., 2.7.12.11., or 2.7.12.12.; or

2. were processed to ensure the destruction of NAI virus (under study), and the necessary precautions were taken after processing to avoid contact of the commodity with any source of NAI virus.

Article 2.7.12.15.

When importing from an NAI free country, zone or compartment, Veterinary Administrations should require:

for poultry semen

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 an NAI free country, zone or compartment for at least the 21 days prior to and at the time of semen collection.

Information concerning the vaccination status of the donor poultry (including the dates of vaccination, and the vaccine used) should be included in the veterinary certificate.

Article 2.7.12.16.

When importing from a HPNAI free country, zone or compartment, Veterinary Administrations should require:

for poultry semen

the presentation of an international veterinary certificate attesting that the donor poultry:

1. came from an HPNAI free country, zone or compartment;
2. were kept in an NAI free establishment for at least 21 days prior to and at the time of semen collection.

1. showed no clinical sign of HPNAI on the day of semen collection;
2. were kept in an HPNAI free country, zone or compartment for at least the 21 days prior to and at the time of semen collection.

Information concerning the vaccination status of the donor flocks (including the dates of vaccination and the vaccine used) should be included in the veterinary certificate.

Article 2.7.12.17.

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Administrations should require:

for semen of birds other than poultry

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 between 7 and 14 days prior to semen collection and shown to be free of NAI infection.

Article 2.7.12.18.

When importing from an NAI free country, zone or compartment, Veterinary Administrations should require:

for fresh meat of poultry
Appendix XVI (contd)

the presentation of an international veterinary certificate attesting that the entire consignment of fresh meat comes from birds:

1. which have been kept in an 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 and have been subjected to ante-mortem and post-mortem inspections for NAI with favourable results.

Article 2.7.12.19.

When importing from a HPNAI free country, zone or compartment, Veterinary Administrations should require:

for fresh meat of poultry
the presentation of an international veterinary certificate attesting that the entire consignment of fresh meat comes from birds:

1. which have been kept in an HPNAI free country, zone or compartment since they were hatched or for at least the past 21 days which have been kept in an establishment since they were hatched or for at least the past 21 days and in which there has been no evidence of NAI in the past 21 days;
2. which have been slaughtered in an approved abattoir and have been subjected to ante-mortem and post-mortem inspections for NAI with favourable results.

Article 2.7.12.20.

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Administrations should require:

for meat products of poultry
the presentation of an international veterinary certificate attesting that:

1. the commodity is derived from fresh meat which meet the requirements of Articles 2.7.12.18. or 2.7.12.19.; or
2. the commodity has been processed to a core temperature of 70ºC for one second (or to an equivalent process) to ensure the destruction of NAI virus (under study);
3. the necessary precautions were taken to avoid contact of the commodity with any source of NAI virus.

Article 2.7.12.21.

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Administrations should require:

for products of poultry origin intended for use in animal feeding, or for agricultural or industrial use
the presentation of an international veterinary certificate attesting that:
1. these commodities come from birds, poultry, which have been kept in an NAI free country, zone or compartment since they were hatched or for at least the past 21 days; or
2. these commodities have been processed to ensure the destruction of NAI virus (under study in accordance with Appendix 3.6.X.);
3. the necessary precautions were taken to avoid contact of the commodity with any source of NAI virus.

Article 2.7.12.22.

Regardless of the NAI status of the country, zone or compartment of origin, Veterinary Administrations should require:

for feathers and down (from poultry)

the presentation of an international veterinary certificate attesting that:

1. these commodities come from birds, poultry, which have been kept in an NAI free country, zone or compartment since they were hatched or for at least the past 21 days; or
2. these commodities have been processed to ensure the destruction of NAI virus (under study);
3. the necessary precautions were taken to avoid contact of the commodity with any source of NAI virus.

Article 2.7.12.23.

Regardless of the NAI status of the country, zone or compartment, Veterinary Administrations should require for the importation of:

meat or other products from birds other than poultry

the presentation of an international veterinary certificate attesting that:

1. the commodity has been processed to ensure the destruction of NAI virus (under study);
2. the necessary precautions were taken after processing to avoid contact of the commodity with any source of NAI virus.

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APPENDIX 3.8.9.
GUIDELINES FOR THE SURVEILLANCE OF AVIAN INFLUENZA

Article 3.8.9.1.

Introduction

This Appendix defines the principles and provides a guide for the surveillance of notifiable avian influenza (NAI) in accordance with Appendix 3.8.1., applicable to countries seeking recognition for a declared NAI status, with or without the use of vaccination. This may be for the entire country, zone or compartment. Guidance for countries seeking free status following an outbreak and for the maintenance of NAI status are provided. This Appendix complements Chapter 2.7.12.

The presence of avian influenza viruses in wild birds creates a particular problem. In essence, no country can declare itself free from avian influenza (AI) in wild birds. However, the definition of NAI in Chapter 2.7.12. refers to the infection in poultry only and this Appendix was 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 guidelines 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 country 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 Member Countries 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 3.8.9.2.

General conditions and methods

1. A surveillance system in accordance with Appendix 3.8.1. should be under the responsibility of the Veterinary Administration. In particular:

   a) a formal and ongoing system for detecting and investigating outbreaks of disease or infection with NAIV 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 Administration. All suspected cases of NAI should be investigated immediately. Where suspicion cannot be resolved by epidemiological and clinical investigation alone, as is frequently the case with low pathogenicity notifiable avian influenza (LPNAI) virus infections, samples should be taken and submitted to an approved 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 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 an 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 sources of NAI.

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 NAI. 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 standstill orders, etc.).

Article 3.8.9.3.

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 NAI 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 country should justify the surveillance strategy chosen as adequate to detect the presence of NAIV infection in accordance with Appendix 3.8.1. and the prevailing epidemiological situation. 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 Country 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 applicant country must justify the choice of design prevalence and confidence level based on the objectives of surveillance and the epidemiological situation, in accordance with Appendix 3.8.1. 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 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 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.
Appendix XVII (contd)

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 until evidence to the contrary is produced.

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 NAI;

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) 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 guidelines 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.

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. The interpretation of serological results in the presence of vaccination is described in 3.8.9.7.

Article 3.8.9.4.

Documentation of NAI or HPNAI free status

1. Countries declaring freedom from NAI or HPNAI for the country, zone or compartment

In addition to the general conditions described in Chapter 2.7.12. of the Terrestrial Code, a Member Country declaring freedom from NAI or highly pathogenic notifiable avian influenza (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 Appendix, 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.
Appendix XVII (contd)

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 3.8.9.5.

Countries, zones or compartments re-declaring regaining freedom from NAI or HPNAI following an outbreak

In addition to the general conditions described in Chapter 2.7.12., a country re-declaring for regaining 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 Country 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 guidelines. The surveillance should at least give the confidence that can be given by a randomized representative sample of the populations at risk.

Article 3.8.9.6.

NAI free establishments within HPNAI free compartments

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 guidelines. The frequency of testing should be based on the risk of infection and at a maximum interval of 21 days.

Article 3.8.9.7.

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 Appendix. 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 ELISA tests. For the HA, antibodies are detected in haemagglutination inhibition (HI) 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.

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 H7N3 in the face of a H7N1 epidemic may be differentiated from infected birds (DIVA) by detection of subtype specific antibodies of the NA protein of the field virus. Alternatively, in the absence of DIVA, inactivated vaccines may induce low titres of antibodies to NP 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. Poultry vaccinated with inactivated whole AI vaccines may develop low titres of antibodies to NSP, but the titre in infected birds will be markedly higher. Alternatively, usage of a vaccine strain with a different NA subtype than the field virus can allow differentiation of vaccinated from infected birds (DIVA) by detection of subtype specific NA antibodies 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 NAI 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:

   **Appendix XVII (contd)**
Appendix XVII (contd)

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)ii) 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.
Figure 1. - Schematic representation of laboratory tests for determining evidence of NAI infection through or following serological surveys

Unvaccinated populations

- NP/M antibodies – ELISA, AGID
  - HI antibodies
    - NAI
    - H1, 4, 6, 8-15
  - S

Vaccinated populations (DIVA)

- Inactivated Whole AIV Vaccine
  - Homologous NA Vaccine
    - Sentinel
    - NSP antibodies
  - Heterologous NA Vaccine
    - Sentinel
    - Heterologous NA antibodies
  - Recombinant vaccine
    - NP/M, NSP or NA antibodies

Virological & Epidemiological Investigation

Serosurveillance

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Figure 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.

Key:

AGID  Agar gel immunodiffusion
DIVA  Differentiating infected from vaccinated animals
ELISA  Enzyme-linked immunosorbant assay
HA  Haemagglutinin
HI  Haemagglutination inhibition
NA  Neuraminidase
NP/M  Nucleoprotein and matrix protein
NSP  Nonstructural protein
S  No evidence of NAIV

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Appendix XVIII

APPENDIX 3.8.X.

GUIDELINES FOR THE INACTIVATION OF THE AVIAN INFLUENZA VIRUS

Article 3.8.X.1.

Egg and egg products

The following industry standard procedures are suitable for the inactivation of highly pathogenic notifiable avian influenza (HPNAI) virus present in egg 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>10% salted yolk</td>
<td>63.3</td>
</tr>
<tr>
<td>Dried egg white</td>
<td>67</td>
</tr>
</tbody>
</table>

Article 3.8.X.2.

Meat

A procedure which produces a core temperature of 70°C for one second is suitable for the inactivation of HPNAI virus present in meat.
APPENDIX 3.2.1.

BOVINE AND SMALL RUMINANT SEMEN

Article 3.2.1.1.

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.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 3.2.1.2.

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 Administration.

3. The centre should be under the supervision and control of the Veterinary Authority 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 a veterinarian designated by the artificial insemination centre and accredited by the Veterinary Administration for relevant official tasks.

Article 3.2.1.3.

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 suspect animals suspected of being infected.
Appendix XIX (contd)

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 OIE-listed ruminant 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.

Article 3.2.1.4.

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 3.2.1.5.

**Conditions applicable to testing of bulls and teaser animals**

Bulls and teaser animals can enter an artificial insemination centre only if they fulfill the following requirements laid down by the Veterinary Administration.

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 2.3.1.5. of the Terrestrial Code.

   b) **Bovine tuberculosis**

      The animals should comply with point 2, 3 or 4 of Article 2.3.3.4. of the Terrestrial Code.

   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.

   d) **Infectious bovine rhinotracheitis-infectious pustular vulvovaginitis (IBR/IPV)**

      If the artificial insemination centre is to be considered as IBR/IPV free, the animals should either:

      i) come from an IBR/IPV free herd as defined in Article 2.3.5.3.; 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 Article 2.2.13.6., 2.2.13.7. or 2.2.13.8., 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 *Trichomonas foetus*, 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)ii) below.)
Appendix XIX (contd)

a) Bovine brucellosis

If the country is not free from 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) Trichomonas 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 Article 2.2.13.9., 2.2.13.10. or 2.2.13.11., 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 ELISA 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 2.3.5.7.

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

Animals 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.

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 Article 2.2.13.9., 2.2.13.10. or 2.2.13.11., 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.
Appendix XIX (contd)

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 2.3.5.3.

Article 3.2.1.6.

Conditions applicable to testing of rams/bucks and teaser animals

Rams/bucks and teaser animals can enter an artificial insemination centre only if they fulfill the following requirements laid down by the Veterinary Administration.

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 2.4.2.6.

b) Ovine epididymitis

The animals should comply with Article 2.4.1.3.

c) Contagious agalactia

The animals should comply with points 1 and 2 of Article 2.4.3.1.

d) Peste des petits ruminants

The animals should comply with points 1, 2, and 4 or 5 of Article 2.4.9.7.

e) Contagious caprine pleuropneumonia

The animals should comply with Article 2.4.6.5. or Article 2.4.6.7., depending on the CCPP status of the country of origin of the animals.

f) Caseous lymphadenitis

The animals should be free from clinical signs for the past 12 months.

g) Paratuberculosis

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

h) Scrapie

If the animals do not originate from a scrapie free country or zone as defined in Article 2.4.8.3., the animals should comply with points 1 and 2 of Article 2.4.8.8.
Appendix XIX (contd)

i) Maedi-visna

The animals should comply with Article 2.4.5.2.

j) Caprine arthritis/encephalitis

The animals should comply with Article 2.4.4.2.

k) Bluetongue

The animals should comply with Article 2.2.13.6., 2.2.13.7. or 2.2.13.8., depending on the bluetongue status of the country of origin of the animals.

l) Tuberculosis

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

m) 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 1 b) or c) of Article 2.4.2.8.

b) Ovine epididymitis

The animals and semen should be subject to testing as described in points 1 d) and 2 of Article 2.4.1.4.

c) Maedi-visna and caprine arthritis/encephalitis or CAE

The animals should be subjected to a serological test.

d) Bluetongue

The animals should comply with the provisions referred to in Article 2.2.13.9., 2.2.13.10. or 2.2.13.11., depending on the bluetongue status of the country of origin of the animals.
Appendix XIX (contd)

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 or CAE;
d) tuberculosis (for goats only);
e) bluetongue.

Article 3.2.1.7.

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 3.2.1.8.

Conditions applicable to the management of bulls, rams and bucks

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.

Measures similar to the above should be adapted to rams and bucks.

Article 3.2.1.9.

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, must be kept clean. A dummy must be cleaned completely after each period of collection. A teaser animal must 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 must 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 must be cleaned completely after each collection. 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 70°C ethyl or 98-99°C isopropyl 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 must 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 3.2.1.10.

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.
   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) 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.
Appendix XIX (contd)

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: either gentamicin (250 µg), tylosin (50 µg), lincomycin-spectinomycin (150/300 µg) or penicillin (500 IU), streptomycin (500 µg), lincomycin-spectinomycin (150/300 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 sterilised with alcohol, ethylene oxide, steam or other approved sterilisation 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 guidelines 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)*.

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. Containers should be sealed with an official numbered seal under the responsibility of the Veterinary Administration before export and accompanied by an international veterinary certificate listing the contents and the number of the official seal.

* 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.

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APPENDIX 3.7.2.

GUIDELINES FOR THE TRANSPORT OF ANIMALS BY SEA

Preamble: These guidelines apply to the following live domesticated animals: cattle, buffalo, deer, camelids, sheep, goats, pigs and equines. They may also be applicable to other domesticated animals.

Article 1

The amount of time animals spend on a journey should be kept to the minimum.

Article 3.7.2.1. bis

Responsibilities

Once the decision to transport the animals by sea has been made, the welfare of the animals during their journey is the paramount consideration and is the joint responsibility of all people involved. These guidelines may also be applied to the transport of animals by water within a country.

The management of animals at post-discharge facilities is outside the scope of this Appendix.

The roles of each of those responsible are defined below:

1. Exporters, owners of animals and managers of facilities are jointly responsible for the general health of the animals and their fitness for the journey, and for their overall welfare during the journey, regardless of whether duties are subcontracted to other parties during transport.

2. The exporter has overall responsibility for the organisation, carrying out and completion of the journey, regardless of whether duties are subcontracted to other parties during transport. The exporter is also responsible for ensuring that equipment and medication are provided as appropriate for the species and journey, and for the presence during the journey of at least one animal handler\(^{48}\) competent for the species being transported. The exporter is also responsible for ensuring compliance of the animals with any required veterinary certification and, in the case of animals for export, any other requirements of the importing and exporting countries.

3. Business or buying/selling agents have a joint responsibility with owners for the selection of animals that are fit to travel. They have a joint responsibility with masters of vessels and managers of facilities at the start and at the end of the journey for the availability of suitable facilities for the assembly, loading, transport, unloading and holding of animals, and for emergencies.

\(^{48}\) An animal handler is a person with a knowledge of the behaviour and needs of animals which, with appropriate experience and a professional and positive response to an animal’s needs, results in effective management and good welfare; their competence should be demonstrated through independent assessment and certification.
Appendix XX (contd)

4. Animal handlers are responsible for the humane handling and care of animals, especially during loading and unloading. To carry out these responsibilities, they should have the authority to take prompt action.

5. The exporter, the shipping company and the master of the vessel are jointly responsible for planning the journey to ensure the care of the animals, including:
   a) choosing appropriate vessels and ensuring that competent animal handlers are available to care for loading and caring for the animals throughout the journey;
   b) developing and keeping up to date contingency plans to address emergencies (including adverse weather conditions) and minimise stress during transport;
   c) correct loading of the ship, regular inspections during the journey and for appropriate responses to problems arising;
   d) disposal of carcasses according to international law.

6. To carry out these responsibilities, the people involved should be competent regarding transport regulations, equipment usage, and the humane handling and care of animals.

7. Managers of facilities during loading of the animals are responsible for:
   a) providing suitable premises for loading the animals;
   b) providing competent animal handlers to load the animals in a manner that causes with minimum stress and injury;
   c) providing appropriate facilities for emergencies;
   d) providing facilities and veterinarians or competent animal handlers capable of killing animals humanely when required.

8. Managers of facilities at the end of the journey are responsible for:
   a) providing suitable facilities for unloading the animals onto transport vehicles for immediate movement or securely holding the animals in lairage, with shelter, water and feed, when required, for transit;
   b) providing competent animal handlers to unload the animals with minimum stress and injury;
   c) minimising the opportunities for disease transmission while the animals are in the facilities;
   d) providing appropriate facilities for emergencies;
   e) providing facilities and veterinarians or competent animal handlers capable of killing animals humanely when required.
9. The responsibilities of the Competent Authority of the exporting country include:
   a) establishing minimum standards for animal welfare, including requirements for inspection of animals before and during their travel, and for certification and record keeping;
   b) approving facilities, containers, vehicles/vessels for the holding and transport of animals;
   c) setting competence standards for animal handlers and managers;
   d) ensuring that the vessel transporting animals meets the required standards, including those of the importing country;
   e) implementation of the standards, including through accreditation of / interaction with other organisations and Competent Authorities;
   f) monitoring and evaluating health and welfare performance, including the use of any veterinary medications.

10. The responsibilities of the Competent Authority of the importing country include:
    a) establishing minimum standards for animal welfare, including requirements for inspection of animals after their travel, and for certification and record keeping;
    b) approving facilities, containers and vehicles for the unloading, holding and transport of animals;
    c) setting competence standards for animal handlers and managers;
    d) implementation of the standards, including through accreditation of / interaction with other organisations and Competent Authorities;
    e) ensuring that the exporting country is aware of the required standards for the vessel transporting the animals;
    f) monitoring and evaluating health and welfare performance, including the use of any veterinary medications.

11. When travelling on vessels with the animals, veterinarians are responsible for the humane handling and treatment of the animals during the journey. To carry out these responsibilities, they should have the authority to act and report independently. The veterinarian should meet with the Master, Chief Officer and the senior animal handler on a daily basis.

12. The receiving Competent Authority should report back to the sending Competent Authority on significant animal welfare problems which occurred during the journey.

Article 3.7.2.2.

Competence

1. All people handling animals or who are otherwise responsible for animals during journeys, should be competent according to their responsibilities listed in Article 3.7.2.1. Competence in areas other than animal welfare would need to be addressed separately. Competence may be gained through formal training and/or practical experience.
Appendix XX (contd)

2. **This** The competence of animal handlers should be demonstrated through a current certificate from an independent body accredited by a Competent Authority. The certificate should be in one of the OIE official languages if the international transport of animals is involved.

3. The assessment of competence for animal handlers should at a minimum address knowledge, and ability to apply that knowledge, in the following areas:
   a) responsibilities for animals during the journey;
   b) sources of advice and assistance;
   c) animal behaviour, general signs of disease, and indicators of poor animal welfare such as stress, pain and fatigue, and their alleviation;
   d) assessment of fitness to travel;
   e) relevant authorities and applicable transport regulations, and associated documentation requirements;
   f) general disease prevention procedures, including cleaning and disinfection;
   g) appropriate methods of animal handling during transport and associated activities such as assembling, loading, and unloading;
   h) methods of inspecting animals, managing situations frequently encountered during transport such as adverse weather conditions, and dealing with emergencies;
   i) species-specific aspects and age-specific aspects of animal handling and care, including feeding, watering and inspection;
   j) appropriate record keeping and maintaining a journey log and other records.

4. Assessment of competence for exporters should at a minimum address knowledge, and ability to apply that knowledge, in the following areas:
   a) planning a journey, including appropriate space allowances, and feed, water and ventilation requirements;
   b) relevant authorities and applicable transport regulations, and associated documentation requirements;
   c) appropriate methods of animal handling during transport and associated activities such as cleaning and disinfection, assembling, loading, and unloading;
   d) species-specific aspects of animal handling and care, including appropriate equipment and medication;
   e) sources of advice and assistance;
   f) appropriate record keeping and maintaining a journey log;
   g) managing situations frequently encountered during transport, such as adverse weather conditions, and dealing with emergencies.
Planning the journey

1. General considerations
   
   a) Adequate planning is a key factor affecting the welfare of animals during a journey.
   
   b) Before the journey starts, plans should be made in relation to:
      
      i) preparation of animals for the journey;
      
      ii) type of transport vessel required;
      
      iii) route, taking into account distance, expected weather and sea conditions;
      
      iv) nature and duration of journey;
      
      v) daily care and management of the animals by providing the appropriate number of animal handlers;
      
      vi) avoiding the mixing of animals from different sources in a single pen group;
      
      vii) provision of appropriate equipment and medication for the numbers and species carried;
      
      viii) emergency response procedures.

2. Preparation of animals for the journey
   
   a) When animals are to be provided with a novel diet e.g. for dry food, and or unfamiliar methods of supplying of feed and or water, they should be preconditioned may be required.
   
   b) There should be planning for water and feed availability during the journey. Feed should be of appropriate quality and composition for the species, age, condition of the animals, etc.
   
   c) Extreme weather conditions are hazards for animals undergoing transport and require appropriate vessel design to minimise risks. Special precautions should be taken for animals that have not been acclimatised or which are unsuited to either hot or cold conditions. In some extreme conditions of heat or cold, animals should not be transported at all.
   
   d) Animals more accustomed to contact with humans and with being handled are likely to be less fearful of being loaded and transported. Animals should be handled and loaded in a manner that reduces their fearfulness and improves their approachability.
   
   e) Behaviour-modifying or other medication should not be used routinely during transport. Such medicines should only be administered when a problem exists in an individual animal, and should be administered by a veterinarian or other person who has been instructed in their use by a veterinarian. Treated animals should be placed in a dedicated area.
Appendix XX (contd)

d) Where there is a potential for spread of infectious disease, and when requested by the Veterinary Authority of the importing country, animals should be vaccinated against diseases to which they are likely to be exposed at their destination.

h) There should be an emergency management plan that identifies the important adverse events that may be encountered during the journey, the procedures for managing each event and the action to be taken in an emergency. For each important event, the plan should document the actions to be undertaken and the responsibilities of all parties involved, including communications and record keeping.

3. Control of disease

As animal transport is often a significant factor in the spread of infectious diseases, journey planning should take into account the following:

a) when possible and agreed by the Veterinary Authority of the importing country, animals should be vaccinated against diseases to which they are likely to be exposed at their destination;

b) medications used prophylactically or therapeutically should only be administered by a veterinarian or other person who has been instructed in their use by a veterinarian;

c) mixing of animals from different sources in a single consignment should be minimized.

4. Vessel and container design and maintenance

a) Vessels used for the sea transport of animals should be designed, constructed and fitted as appropriate to the species, size and weight of the animals to be transported. Special attention should be paid to the avoidance of injury to animals through the use of secure smooth fittings free from sharp protrusions and the provision of non-slip flooring. The avoidance of injury to animal handlers while carrying out their responsibilities should be emphasised.

b) Vessels should be designed to permit thorough cleaning and disinfection, and the management of faeces and urine.

c) Vessels and their fittings should be maintained in good mechanical and structural condition.

d) Vessels should have adequate ventilation to meet variations in climate and the thermo-regulatory needs of the animal species being transported. The ventilation system should be capable of operating effective when the vessel is stationary and the air flow should be adjustable. An emergency power supply should be available to maintain ventilation in the case of primary machinery breakdown.

e) The feeding and watering system should be designed to permit adequate access to feed and water appropriate to the species, size and weight of the animals, and to minimise soiling of pens.

f) Vessels should be designed so that the faeces or urine from animals on upper levels do not soil animals on lower levels, or their feed or water.
g) Loading and stowage of feed and bedding should be carried out in such a way to ensure protection from fire hazards, the elements and sea water.

h) Where appropriate, suitable bedding, such as straw or sawdust, should be added to vessel floors to assist absorption of urine and faeces, provide better footing for animals and protect animals (especially young animals) from hard or rough flooring surfaces and adverse weather conditions.

i) The above principles apply also to containers used for the transport of animals.

5. Special provisions for transport in road vehicles on roll-on/roll-off vessels or for containers

a) Road vehicles and containers should be equipped with a sufficient number of adequately designed, positioned and maintained securing points enabling them to be securely fastened to the vessel.

b) Road vehicles and containers should be secured to the ship before the start of the sea journey to prevent them being displaced by the motion of the vessel.

c) Vessels should have adequate ventilation to meet variations in climate and the thermo-regulatory needs of the animal species being transported, especially where the animals are transported in a secondary vehicle/container on enclosed decks.

d) Due to the risk of limited airflow on certain vessels’ decks, a road vehicle or container may require a forced ventilation system of greater capacity than that provided by natural ventilation.

6) Nature and duration of the journey

The maximum duration of a journey should be determined according to:

a) the ability of the animals to cope with the stress of transport (such as very young, old, lactating or pregnant animals);

b) the animals’ previous transport experience;

c) the likely onset of fatigue;

d) the need for special attention;

e) the need for feed and water;

f) the increased susceptibility to injury and disease;


g) space allowance and vessel design;

h) weather conditions.

7. Space allowance

a) The number of animals which should be transported on a vessel and their allocation to different pens on the vessel should be determined before loading.
Appendix XX (contd)

b) The amount of space required, including headroom, depends on the species of animal and should allow the necessary thermoregulation. Each animal should be able to assume its natural position for transport (including during loading and unloading) without coming into contact with the roof or upper deck of the vessel. When animals lie down, there should be enough space for every animal to adopt a comfortable, normal lying posture.

c) Calculations for the space allowance for each animal should be carried out, using the figures given in these guidelines Appendix XXX or, in their absence, in a relevant national or international document. The size of pens will affect the number of animals in each.

d) The same principles apply when animals are transported in containers.

8. Ability to observe animals en route during the journey

a) Animals should be positioned to enable them to be observed regularly and clearly by the animal handler or other responsible person, during the journey to ensure their safety and good welfare.

b) To allow an adequate inspection of animals en route, it should be possible for each animal to be clearly observed by the animal handler or other responsible person.

9. Emergency response procedures

Appropriate contingency plans to address emergencies should be prepared in advance.

There should be an emergency management plan that identifies the important adverse events that may be encountered during the journey, the procedures for managing each event and the action to be taken in an emergency. For each important event, the plan should document the actions to be undertaken and the responsibilities of all parties involved, including communications and record keeping.

Article 3.7.2.4.

Documentation

1. Animals should not be loaded until the documentation required to that point is complete.

2. The documentation accompanying the consignment should include:
   a) journey travel plan (including an emergency management plan);
   b) time, date and place of loading;
   c) the journey log – a daily record of inspection and important events which includes records of morbidity and mortality and actions taken, climatic conditions, food and water consumed, medication provided, mechanical defects;
   d) expected time, date and place of arrival and unloading;
   e) veterinary certification, when required;
Appendix XX (contd)

f) animal identification to allow traceback of individual animals to the premises of departure, and, where possible, to the premises of origin;

g) details of any animals considered ‘at risk’ (Article 3.7.2.5);

h) number of animal handlers on board, and their competencies;

i) stocking density estimate for each load in the consignment.

3. When veterinary certification should be required to accompany consignments of animals and, it should address:

a) when required, cleaning and details of disinfection carried out of the vessel;

b) fitness of the animals to travel;

c) animal identification (description, number, etc.);

d) health status including any tests, treatments and vaccinations carried out, if required.

Article 3.7.2.5.

Pre-journey period

1. General considerations

a) Before each journey, vessels should be thoroughly cleaned and, if necessary, treated for animal and public health purposes, using chemicals approved by the Competent Authority. When cleaning is necessary during a journey, this should be carried out with the minimum of stress to the animals.

b) In some circumstances, animals may require pre-journey assembly. In these circumstances, the following points should be considered:

i) Pre-journey rest is necessary if the welfare of animals has become poor during the collection period because of the physical environment or the social behaviour of the animals.

ii) For animals such as pigs which are susceptible to motion sickness, and in order to reduce urine and faeces production during the journey, a short period of feed deprivation prior to loading is desirable.

iii) When animals will be provided with a novel diet or method of water provision during or after transport, an adequate period of pre-exposure is necessary. Preconditioning to the feed to be used on the vessel may be necessary in such cases.

iv) When animals are to be provided with a novel diet or unfamiliar methods of supplying of feed or water, they should be preconditioned.
Appendix XX (contd)

c) Where an animal handler believes that there is a significant risk of disease among the animals to be loaded or significant doubt as to their fitness to travel, the animals should be examined by a veterinarian.

d) Pre-journey assembly /holding areas should be designed to:

i) securely contain the animals;

ii) maintain an environment safe from hazards, including predators and disease;

iii) protect animals from exposure to adverse weather conditions; and

iv) allow for maintenance of social groups; and

v) allow for rest, watering and feeding.

2. Selection of compatible groups

Compatible groups should be selected before transport to avoid adverse animal welfare consequences. The following guidelines should be applied when assembling groups of animals:

a) animals of different species should not be mixed unless they are judged to be compatible;

b) animals of the same species can be mixed unless there is a significant likelihood of aggression; aggressive individuals should be segregated (recommendations for specific species are described in detail in Article 3.7.2.10.). For some species, animals from different groups should not be mixed because poor welfare occurs unless they have established a social structure;

c) young or small animals may need to be separated from older or larger animals, with the exception of nursing mothers with young at foot;

d) animals with horns or antlers should not be mixed with animals lacking horns or antlers, unless judged to be compatible;

e) animals reared together should be maintained as a group; animals with a strong social bond, such as a dam and offspring, should be transported together.

3. Fitness to travel

a) Animals should be inspected by a veterinarian or an animal handler to assess fitness to travel. If its fitness to travel is in doubt, the animal should be examined by a veterinarian. Animals found unfit to travel before travel and those found unfit to travel by farm staff, an animal handler or a veterinarian, should not be loaded onto a vessel.

b) Humane and effective arrangements should be made by the owner or agent for the handling and care of any animal rejected as unfit to travel.
Appendix XX (contd)

c) Animals that are unfit to travel include:
   i) those that are sick, injured, weak, disabled or fatigued;
   ii) those that are unable to stand unaided and bear weight on each leg;
   iii) those that are blind in both eyes;
   iv) those that cannot be moved without causing them additional suffering;
   v) newborn with an unhealed navel;
   vi) females travelling without young which have given birth within the previous 48 hours;
   vii) pregnant animals which would be in the final 10% of their gestation period at the planned time of unloading.

d) Risks during transport can be reduced by selecting animals best suited to the conditions of travel and those that are acclimatised to expected weather conditions.

e) Animals at risk, and requiring better conditions and additional attention during transport include:
   i) very large or obese individuals;
   ii) very young or old animals;
   iii) excitable or aggressive animals;
   iv) animals subject to motion sickness;
   v) animals which have had little contact with humans;
   vi) females in the last third of pregnancy or in heavy lactation.

f) Hair or wool length needs consideration should be considered in relation to the weather conditions expected during transport.

Article 3.7.2.6.

Loading

1. Experienced Competent supervision
   a) Loading should be carefully planned as it has the potential to be the cause of poor welfare in transported animals.
   
   b) Loading should be supervised by the Competent Authority and managed conducted by an animal handler(s). Animal handlers should ensure that animals are loaded quietly and without unnecessary noise, harassment or force, and that untrained assistants or spectators do not impede the process.
Appendix XX (contd)

e) Ventilation during loading and the journey should provide for fresh air, and the removal of excessive heat, humidity and noxious fumes (such as ammonia and carbon monoxide). Under warm and hot conditions, ventilation should allow for the adequate convective cooling of each animal. In some instances, adequate ventilation can be achieved by increasing the space allowance for animals.

2. Facilities

a) The facilities for loading including the collecting area at the wharf, races and loading ramps should be designed and constructed to take into account of the needs and abilities of the animals with regard to dimensions, slopes, surfaces, absence of sharp projections, flooring, sides, etc.

b) Ventilation during loading and the journey should provide for fresh air, and the removal of excessive heat, humidity and noxious fumes (such as ammonia and carbon monoxide). Under warm and hot conditions, ventilation should allow for the adequate convective cooling of each animal. In some instances, adequate ventilation can be achieved by increasing the space allowance for animals.

c) All loading facilities should be properly illuminated to allow the animals to be easily inspected by the animal handler(s), and to allow the animals’ ease of movement at all times. Facilities should provide uniform lighting light levels directly over approaches to sorting pens, chutes, loading ramps, with brighter lighting light levels inside vehicles / containers, in order to minimise baulking. Dim lighting light levels may be advantageous for the catching of some animals. Artificial lightening may be required.

3. Goads and other aids

The following principles should apply:

a) Goads (aids for encouraging animals to move) should not be used on Animals that have little or no room to move should not be subjected to physical force or goads and other aids which compel movement.

b) Useful and permitted goads include panels, flags, plastic paddles, flappers (a length of cane with a short strap of leather or canvas attached), plastic bags and metallic rattles; they should be used in a manner sufficient to encourage and direct movement of the animals but without physical contact with them.

c) Painful procedures (including whipping, tail twisting, use of nose twitches, pressure on eyes, ears or external genitalia), or the use of unsuitable goads or other aids (including sticks with sharp ends, lengths of metal piping, fencing wire or heavy leather belts), should not be used to move animals.

d) Unsuitable goads such as large wooden sticks, sticks with sharp ends, lengths of metal piping, fencing wire or heavy leather belts should not be used to strike animals.
Appendix XX (contd)

d) The use of goads which administer electric shocks should be discouraged, and restricted to that necessary to assist movement of the animal. If such use is necessary, it should be limited to battery-powered goads on the hindquarters of pigs and large ruminants, and never on sensitive areas such as the eyes, mouth, ears, anogenital region or belly. Such instruments should not be used on horses, sheep and goats of any age, or on calves or piglets.

e) Shouting or yelling at animals or making loud noises eg through the cracking of whips to encourage them to move should not occur, as such actions may make the animals agitated, leading to crowding or falling.

f) The use of well trained dogs to help with the loading of some species may be acceptable.

g) Manual lifting is permissible for young animals that may have difficulty negotiating ramps, but the lifting of animals by body parts such as their tail, head, horns, ears, limbs, wool or hair should not be permitted. The throwing or dropping of animals should not be permitted.

Article 3.7.2.7.

Travel

1. General considerations

   a) Animal handler(s) should check the consignment immediately before departure to ensure that the animals have been loaded according to the load plan. Each consignment should be checked again within 24 hours.

   b) Adjustments should be made to the stocking density within 48 hours of departure and as appropriate during the journey.

   c) Each pen of animals should be observed on a daily basis for normal behaviour, health and welfare, and the correct operation of ventilation, watering and feeding systems. There should also be a night patrol. Any necessary corrective action should be undertaken promptly.

   d) Adequate access to suitable feed and water should be ensured for all animals in each pen.

2. Sick and injured animals

   a) Sick and injured animals should be segregated/isolated if possible.

   b) Sick and injured animals should be appropriately treated promptly and or humanely killed, in accordance with a predetermined emergency response plan (Article 3.7.2.3), and Veterinary advice should be sought if necessary. All drugs and products should be used in accordance with the manufacturer’s or veterinarian’s recommendations.

   c) A record of treatments carried out and their outcomes should be kept.
Appendix XX (contd)

d) When euthanasia is necessary, the person responsible for the animals must ensure that it is carried out humanely, and results in immediate death. When necessary, Assistance should be sought from a veterinarian or other person(s) competent in euthanasia procedures. Recommendations for specific species are described in Appendix 3.7.6. on humane killing of animals for disease control purposes.

3. Cleaning and disinfection

a) Vessels and containers used to carry the animals should be cleaned before re-use through the physical removal of manure and bedding by scraping, washing and flushing vessels and containers with water. This should be followed by disinfection when there are concerns about disease transmission.

b) Manure, litter and bedding should be disposed of in such a way as to prevent the transmission of disease and in compliance with all relevant health and environmental legislation.

c) Where cleaning or disinfection is necessary during travel, it should be carried out with the minimum stress to the animals.

Article 3.7.2.8.

Unloading and post-journey handling

1. General considerations

a) The required facilities and the principles of animal handling detailed in Article 3.7.2.6. apply equally to unloading, but consideration should be given to the likelihood that the animals will be fatigued.

b) Unloading should be carefully planned as it has the potential to be the cause of poor welfare in transported animals.

c) A livestock vessel should have priority attention when arriving in port and have priority access to a berth with suitable unloading facilities. As soon as possible after the ship’s arrival at the port and acceptance of the consignment by the Competent Authority, animals should be unloaded into appropriate facilities.

d) The accompanying veterinary certificate and other documents should meet the requirements of the importing country. Veterinary inspections should be completed as quickly as possible.

e) Unloading should be supervised by the Competent Authority and managed conducted by an competent animal handler(s). The animal handlers should ensure that animals are unloaded as soon as possible after arrival but sufficient time should be allowed for unloading to proceed quietly and without unnecessary noise, harassment or force, and that untrained assistants or spectators do not impede the process.
2. Facilities
   a) The facilities for unloading including the collecting area at the wharf, races and unloading ramps should be designed and constructed to take into account of the needs and abilities of the animals with regard to dimensions, slopes, surfaces, absence of sharp projections, flooring, sides, etc.

   b) All unloading facilities should be properly illuminated, have sufficient lighting to allow the animals to be easily inspected by the animal handler(s), and to allow the animals’ ease of movement at all times.

   c) In case of emergencies, Port facilities should provide animals with appropriate care and comfort, adequate space, access to quality feed and clean drinking water, and shelter from extreme weather conditions.

3. Sick and injured animals
   a) An animal that has become sick, injured or disabled during a journey should be appropriately treated or humanely killed (see Appendix 3.7.6.). When necessary, veterinary advice should be sought in the care and treatment of these animals.

   b) In some cases, where animals are non-ambulatory due to fatigue, injury or sickness, it may be in the best welfare interests of the animal to be treated or euthanased aboard the vessel.

   c) If unloading is in the best welfare interests of animals that are fatigued, injured or sick, there should be appropriate facilities and equipment for the humane unloading of such animals. These animals should be unloaded in a manner that causes the least amount of suffering. After unloading, separate pens and other appropriate facilities and treatments should be provided for sick or injured animals.

4. Cleaning and disinfection
   a) Vessels and containers used to carry the animals should be cleaned before re-use through the physical removal of manure and bedding, by scraping, washing and flushing vessels and containers with water until visibly clean. This should be followed by disinfection when there are concerns about disease transmission.

   b) Manure, litter and bedding should be disposed of in such a way as to prevent the transmission of disease and in compliance with all relevant health and environmental legislation.

   c) Where cleaning or disinfection is necessary during travel, it should be carried out with the minimum of stress to the animals.

Article 3.7.2.9.

Actions in the event of a refusal to allow the importation of a shipment

1. The welfare of the animals should be the first consideration in the event of a refusal to import.
Appendix XX (contd)

2. When a shipment has been refused import, the Competent Authority of that country should make available suitable isolation facilities to allow the unloading of animals from a vessel and their secure holding, without posing a risk to the health of the national herd, pending resolution of the situation. In this situation, the priorities should be:
   a) the Competent Authority of the importing country should provide urgently in writing the reasons for the refusal;
   b) in the event of a refusal for animal health reasons, the Competent Authority of the importing country should provide urgent access to an OIE-appointed veterinarian(s) to assess the animals’ health status with regard to the importing country’s concerns, and the necessary facilities and approvals to expedite the required diagnostic testing;
   c) the Competent Authority of the importing country should provide access to allow continued assessment of the ongoing health and welfare situation;
   d) if the matter cannot be promptly resolved, the Competent Authority of the exporting and importing countries should call on the OIE to mediate.

3. In the event that the animals are required to remain on the vessel, the priorities should be:
   a) the Competent Authority of the importing country should allow rep rovision of the vessel with water and feed as necessary;
   b) the Competent Authority of the importing country should provide urgently in writing the reasons for the refusal;
   c) in the event of a refusal for animal health reasons, the Competent Authority of the importing country should provide urgent access to an OIE-appointed veterinarian(s) to assess the animals’ health status with regard to the importing country’s concerns, and the necessary facilities and approvals to expedite the required diagnostic testing;
   d) the Competent Authority of the importing country should provide access to allow continued assessment of the ongoing health and welfare situation other aspects of the welfare of the animals, and the necessary actions to deal with any issues which arise;
   e) if the matter cannot be urgently resolved, the Competent Authorities of the exporting and importing countries should call on the OIE to mediate.

4. The OIE should utilise its dispute settlement mechanism to identify a mutually agreed solution which will address the animal health and welfare issues in a timely manner.

Article 3.7.2.10.

Species specific issues

Cattle are sociable animals and may become agitated if they are singled out. Social order is usually established at about two years of age. When groups are mixed, social order has to be re-established and aggression may occur until a new order is established. Crowding of cattle may also increase aggression as the animals try to maintain personal space. Social behaviour varies with age, breed and sex; Bos indicus and B. indicus-cross animals are usually more temperamental than European breeds. Young bulls, when moved in groups, show a degree of playfulness (pushing and shoving) but become more aggressive and territorial with age. Adult bulls have a minimum personal space of six square metres. Cows with young calves can be very protective, and handling calves in the presence of their mothers can be dangerous.
Appendix XX (contd)

**Goats** should be handled calmly and are more easily led or driven than if they are excited. When goats are moved, their gregarious tendencies should be exploited. Activities which frighten, injure or cause agitation to animals should be avoided. Bullying is particularly serious in goats. Housing strange goats together could result in fatalities, either through physical violence, or subordinate goats being refused access to food and water.

**Sheep** are sociable animals with good eyesight and tend to “flock together”, especially when they are agitated. They should be handled calmly and their tendency to follow each other should be exploited when they are being moved. Sheep may become agitated if they are singled out for attention and will strive to rejoin the group. Activities which frighten, injure or cause agitation to sheep should be avoided. They can negotiate steep ramps.

**Pigs** have poor eyesight, and may move reluctantly in strange surroundings. They benefit from well lit loading bays. Since they negotiate ramps with difficulty, these should be as level as possible and provided with secure footholds. Ideally, a hydraulic lift should be used for greater heights. Pigs also negotiate steps with difficulty. A good ‘rule-of-thumb’ is that no step should be higher than the pig’s front knee. Serious aggression may result if unfamiliar animals are mixed. Pigs are highly susceptible to heat stress.

**Horses** in this context include all solipeds, donkeys, mules, hinnies and zebra. They have good eyesight and a very wide angle of vision. They may have a history of loading resulting in good or bad experiences. Good training should result in easier loading, but some horses can prove difficult, especially if they are inexperienced or have associated loading with poor transport conditions. In these circumstances, two experienced handlers can load an animal by linking arms or using a strop below its rump. Blindfolding may even be considered. Ramps should be as shallow as possible. Steps are not usually a problem when horses mount a ramp, but they tend to jump a step when descending, so steps should be as low as possible. Horses benefit from being individually stalled, but may be transported in compatible groups. When horses are to travel in groups, their shoes should be removed.

**Camelids** in this context comprise llamas, alpacas, guanaco and vicuna. They have good eyesight and, like sheep, can negotiate steep slopes, though ramps should be as shallow as possible. They load most easily in a bunch as a single animal will strive to rejoin the others. Whilst they are usually docile, they have an unnerving habit of spitting in self-defence. During transport, they usually lie down. They frequently extend their front legs forward when lying, so gaps below partitions should be high enough so that their legs are not trapped when the animals rise.

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*OIE Terrestrial Animal Health Standards Commission/September 2005*
Preamble: These guidelines apply to the following live domesticated animals: cattle, buffalo, camels, sheep, goats, pigs, poultry and equines. They will also be largely applicable to some other animals (e.g. deer, other camelids and ratites). Wild, feral and partly domesticated animals may need different conditions.

Article 1

The amount of time animals spend on a journey should be kept to the minimum.

Article 3.7.3.1. bis

Responsibilities

Once the decision to transport the animals has been made, the welfare of the animals during their journey is the paramount consideration and is the joint responsibility of all people involved.

The roles of each of those responsible are defined below:

1. The owners and managers of the animals are responsible for the general health of the animals and their fitness for the journey, and for their overall welfare during the journey, regardless of whether duties are subcontracted to other parties during transport. They are also responsible for ensuring compliance with any required veterinary or other certification, and for the presence during the journey of at least one animal handler\(^49\) competent for the species being transported, with the authority to take prompt action. They are also responsible for ensuring that equipment and veterinary assistance are provided as appropriate for the species and journey.

2. Business agents or buying/selling agents have a joint responsibility with owners for the selection of animals that are fit to travel. They have a joint responsibility with market owners and managers of facilities at the start and at the end of the journey for the availability of suitable facilities for the assembly, loading, transport, unloading and holding of animals, and for emergencies.

3. Animal handlers are responsible for the humane handling and care of the animals, especially during loading and unloading, and for maintaining a journey log. To carry out their responsibilities, they should have the authority to take prompt action. In the absence of a separate animal handler, the driver is the animal handler.

4. Transport companies, vehicle owners and drivers are responsible for planning the journey to ensure the care of the animals:
   a) transport companies and vehicle owners are responsible for choosing appropriate vehicles and ensuring that properly trained staff are available for loading and caring for animals;

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\(^49\) An animal handler is a person with a knowledge of the behaviour and needs of animals which, with appropriate experience and a professional and positive response to an animal's needs, results in effective management and good welfare; their competence should be demonstrated through independent assessment and certification.
Appendix XXI (contd)

b) transport companies and vehicle owners are responsible for developing and keeping up to date contingency plans to address emergencies and minimise stress during transport;

c) transport companies and vehicle owners are responsible for producing a journey plan which includes a loading plan, journey duration and location of resting places;

d) drivers are responsible for loading only those animals which are fit to travel, for their correct loading into the vehicle and their inspection during the journey, and for appropriate responses to problems arising. If its fitness to travel is in doubt, the animal should be examined by a veterinarian in accordance with point 5 a) of Article 3.7.3.5.

5. Managers of facilities at the start and at the end of the journey and at resting points are responsible for:

a) providing suitable premises for loading, unloading and securely holding the animals, with water and feed when required, until further transport, sale or other use (including rearing or slaughter);

b) providing competent animal handlers to load, unload, drive and hold animals in a manner that causes minimum stress and injury;

c) minimising the opportunities for disease transmission;

d) providing appropriate facilities, with water and feed when required;

e) providing appropriate facilities for emergencies;

f) providing facilities for washing and disinfecting vehicles after unloading;

g) providing facilities and competent staff to allow the humane killing of animals when required;

b) ensuring proper rest times and minimal delay during stops.

6. The responsibilities of Competent Authorities include:

a) establishing minimum standards for animal welfare, including requirements for inspection of animals before, during and after their travel, and appropriate certification and record keeping;

b) approving facilities, containers and vehicles for the transport of animals;

c) setting standards for the competence of drivers, animal handlers and managers;

d) ensuring appropriate awareness and training of drivers, animal handlers and managers;

e) implementation of the standards, including through accreditation of / interaction with other organisations;

f) monitoring and evaluating the effectiveness of standards of health and other aspects of welfare;

g) monitoring and evaluating the use of veterinary medications.
7. All individuals, including veterinarians, involved in transporting animals and the associated handling procedures should receive appropriate training and be competent to meet their responsibilities.

8. The receiving Competent Authority should report back to the sending Competent Authority on significant animal welfare problems which occurred during the journey.

Article 3.7.3.2.

Competence

1. All people handling animals, or who are otherwise responsible for animals during journeys, should be competent according to their responsibilities listed in Article 3.7.3.1. Competence may be gained through formal training and/or practical experience. Competence in areas other than animal welfare would need to be addressed separately.

2. The competence of animal handlers should be demonstrated through a current certificate from an independent body, accredited by the Competent Authority. The certificate should be in one of the OIE official languages if the international transport of animals is involved.

3. The assessment of the competence of animal handlers should at a minimum address knowledge, and ability to apply that knowledge, in the following areas:
   a) planning a journey, including appropriate space allowance, and feed, water and ventilation requirements;
   b) responsibilities for animals during the journey, including loading and unloading;
   c) sources of advice and assistance;
   d) animal behaviour, general signs of disease, and indicators of poor animal welfare such as stress, pain and fatigue, and their alleviation;
   e) assessment of fitness to travel;
   f) relevant authorities and applicable transport regulations, and associated documentation requirements;
   g) general disease prevention procedures, including cleaning and disinfection;
   h) appropriate methods of driving;
   i) methods of inspecting animals, managing situations frequently encountered during transport such as adverse weather conditions, and dealing with emergencies;
   j) species-specific and age-specific aspects of animal handling and care, including feeding, watering and inspection;
   k) maintaining a journey log and other records.
Appendix XXI (contd)

Article 3.7.3.3.

Planning the journey

1. General considerations

   a) Adequate planning is a key factor affecting the welfare of animals during a journey.

   b) Before the journey starts, plans should be made in relation to:

      i) preparation of animals for the journey;
      ii) choice of road or rail;
      iii) nature and duration of the journey;
      iv) vehicle / container design and maintenance, including roll-on roll-off vessels;
      v) required documentation;
      vi) space allowance;
      vii) rest, water and feed;
      viii) observation of animals en route;
      ix) control of disease; and
      x) emergency response procedures.

   c) Regulations concerning drivers (for example, maximum driving periods) should be harmonised with maximum transport journey intervals appropriate for the species.

2. Preparation of animals for the journey

   a) When animals are to be provided with a novel diet or method of water provision during transport, an adequate period of adaptation should be planned. For animals such as pigs which are susceptible to motion sickness, and in order to reduce urine and faeces production during the journey, a short period of feed deprivation prior to loading may be desirable.

   b) Animals should be exposed to appropriate contact with humans and handling conditions (including methods of restraint) prior to transport to reduce their fearfulness and improve their approachability (see Article 3.7.3.5.). Since animals more accustomed to contact with humans and with being handled are likely to be less fearful of being loaded and transported. People handling animals should handle and load animals in a manner that reduces their fearfulness and improves their approachability.

   c) Behaviour-modifying compounds (such as tranquillisers) should not be used routinely during transport. Such compounds should only be administered when a problem exists in an individual animal, and should be administered by a veterinarian or other person who has been instructed in their use by a veterinarian.
3. **Nature and duration of the journey**

The maximum duration of a journey should be determined according to:

a) the ability of the animals to cope with the stress of transport (such as very young, old, lactating or pregnant animals);

b) the animals’ previous transport experience;

c) the onset of fatigue;

d) the need for special attention;

e) the need for feed and water;

f) the increased susceptibility to injury and disease;

g) space allowance, vehicle design, road conditions and driving quality;

h) weather conditions.

4. **Vehicle and container design and maintenance**

a) Vehicles and containers used for the transport of animals should be designed, constructed and fitted as appropriate to the species, size and weight of the animals to be transported; special attention should be paid to the avoidance of injury to animals through the use of secure smooth fittings free from sharp protrusions. The avoidance of injury to drivers and animal handlers while carrying out their responsibilities should be emphasised.

b) Vehicles and containers should be designed with the structures necessary to provide protection from adverse weather conditions and to minimise the opportunity for animals to escape.

c) In order to minimise the likelihood of the spread of pathogenic agents, infectious disease during transport, vehicles and containers should be designed to permit thorough cleaning and disinfection, and the containment of faeces and urine during a journey.

d) Vehicles and containers should be maintained in good mechanical and structural condition.

e) Vehicles and containers should have adequate ventilation to meet variations in climate and the thermo-regulatory needs of the animal species being transported; the ventilation system should be capable of operating effectively when the vehicles is stationary and the air flow should be adjustable.

f) Vehicles should be designed so that the faeces or urine from animals on upper levels do not soil animals on lower levels, nor their feed and water.

g) When vehicles are carried on board ferries, facilities for adequately securing them should be available.
Appendix XXI (contd)

h) If feeding or watering while the vehicle is moving is required, adequate facilities on the vehicle should be available.

i) When appropriate, suitable bedding should be added to vehicle floors to assist absorption of urine and faeces, to minimise slipping by animals, and protect animals (especially young animals) from hard flooring surfaces and adverse weather conditions.

5. Special provisions for transport in vehicles (road and rail) on roll-on/roll-off vessels or for containers

a) Vehicles and containers should be equipped with a sufficient number of adequately designed, positioned and maintained securing points enabling them to be securely fastened to the vessel.

b) Vehicles and containers should be secured to the ship before the start of the sea journey to prevent them being displaced by the motion of the vessel.

c) Roll-on/roll-off vessels should have adequate ventilation to meet variations in climate and the thermo-regulatory needs of the animal species being transported, especially where the animals are transported in a secondary vehicle/container on enclosed decks.

6. Space allowance

a) The number of animals which should be transported on a vehicle or in a container and their allocation to different compartments should be determined before the vehicle or container is loaded.

b) The space required on a vehicle or in a container depends upon whether or not the animals need to lie down (for example, pigs, camels and poultry), or to stand (horses). Animals which will need to lie down often stand when first loaded or when the vehicle is driven with too much lateral movement or sudden braking.

c) When animals lie down, they should all be able to adopt a comfortable, normal lying posture which allows necessary thermoregulation.

d) When animals are standing, they should have sufficient space to adopt a balanced position as appropriate to the climate and species transported (Article XXX).

e) The amount of headroom necessary depends on the species of animal. Each animal should be able to assume its natural position for transport (including during loading and unloading) without coming into contact with the roof or upper deck of the vehicle.

f) Calculations according to for the space allowance permitted for each animal should be carried out using the figures given in Appendix XXX or, in their absence, in a relevant national or international document. The size of already established groups will affect the number and size of the pens, and the distribution of animals in pens on the vehicle. The number and size of pens on the vehicle should be varied to where possible accommodate already established groups of animals while avoiding group sizes which are too large.
g) Other factors which may influence space allowance include:
   i) vehicle / container design;
   ii) length of journey;
   iii) need to provide feed and water on the vehicle;
   iv) quality of roads;
   v) expected weather conditions.

7. Rest, water and feed
   a) There should be planning for the availability of suitable water and feed during the journey. Feed should be of appropriate quality and composition for the species, age, condition of the animals, climatic conditions, etc. as appropriate and needed for the species, age, and condition of the animals, as well as the duration of the journey, climatic conditions, etc.
   b) Animals should be rested. There should be planning for the resting of animals at resting points at appropriate intervals during the journey. The type of transport, the age and species of the animals being transported should determine the frequency of rest stops and whether the animals are should be unloaded. There should be planning for water and feed availability during rest stops.

8. Ability to observe animals en route in relation to during the journey duration
   a) Animals should be positioned to enable each animal to be observed regularly during the journey to ensure their safety and good welfare.
   b) If the animals are in crates or on multi-tiered vehicles which do not allow free access for observation, for example where the roof of the tier is too low (i.e. less than 1.3 m), animals cannot be inspected adequately, and serious injury or disease could go undetected. In these circumstances, a shorter journey duration should be allowed, and the maximum duration will vary according to the rate at which problems arise in the species and under the conditions of transport.

9. Control of disease
   As animal transport is often a significant factor in the spread of infectious diseases, journey planning should take the following into account:
   a) mixing of animals from different sources in a single consignment should be minimised;
   b) contact at resting points between animals from different sources should be avoided;
   c) when possible, animals should be vaccinated against diseases to which they are likely to be exposed at their destination;
Appendix XXI (contd)

d) medications used prophylactically or therapeutically should only be administered by a veterinarian or other person who has been instructed in their use by a veterinarian and agreed by the Veterinary Authority of the importing country.

10. Emergency response procedures

Appropriate contingency plans to address emergencies should be prepared in advance.

There should be an emergency management plan that identifies the important adverse events that may be encountered during the journey, the procedures for managing each event and the action to be taken in an emergency. For each important event, the plan should document the actions to be undertaken and the responsibilities of all parties involved, including communications and record keeping.

11. Other considerations

a) Extreme weather conditions are hazardous for animals undergoing transport and require appropriate vehicle design to minimise risks. Special precautions should be taken for animals that have not been acclimatised or which are unsuited to either hot or cold conditions. In some extreme conditions of heat or cold, animals should not be transported at all.

b) In some circumstances, transportation during the night may reduce thermal stress or the adverse effects of other external stimuli.

Article 3.7.3.4.

Documentation

1. Animals should not be loaded until the required documentation required to that point is complete.

2. The documentation accompanying the consignment should include:

   a) journey travel plan (including an emergency management plan);
   b) date, time, and place of loading and unloading;
   c) veterinary certification, when required;
   d) driver’s competencies;
   e) identities of the animals transported to allow traceback of individual animals to the premises of departure and, where possible, to the premises of origin;
   f) details of any animals considered ‘at risk’ (Article 3.7.3.5.);
   g) documentation of the period of rest, and access to feed and water, prior to the journey;
   h) stocking density estimate for each load in the consignment;
Appendix XXI (contd)

i) the journey log - daily record of inspection and important events, including records of morbidity and mortality, actions taken, climatic conditions, rest stops, travel time and distance, feed and water offered and estimates of consumption, medication provided, and mechanical defects.

3. When veterinary certification is required to accompany consignments of animals, it should include:

a) fitness of animals to travel;

b) appropriate animal identification (description, number, etc.);

c) health status including any tests, treatments and vaccinations status carried out;

d) when required, details of disinfection carried out.

At the time of certification, the veterinarian should notify the animal handler of any factors affecting the animals’ fitness to travel for a particular journey.

Article 3.7.3.5.

Pre-journey period

1. General considerations

a) Pre-journey rest is necessary if the welfare of animals has become poor during the collection period because of the physical environment or the social behaviour of the animals.

b) Pre-journey assembly/holding areas should be designed to:

   i) securely hold the animals;

   ii) maintain a safe environment from hazards, including predators and disease;

   iii) protect animals from exposure to severe weather conditions;

   iv) allow for maintenance of social groups, and

   v) allow for rest, and appropriate water and feed.

c) Consideration should be given to an animal's previous transport experience, training and conditioning if known as these may reduce fear and stress in animals.

d) Feed and water should be provided pre-journey if the journey duration is greater than the normal inter-feeding and drinking interval for the animal. Recommendations for specific species are described in detail in Article 3.7.3.10.

e) When animals will be provided with a novel diet or method of feed or water provision during or after transport, an adequate period of adaptation should be planned; pre-exposure is necessary.
Appendix XXI (contd)

f) Before each journey, vehicles and containers should be thoroughly cleaned and, if necessary, treated for animal health and public health purposes, using methods approved by the Competent Authority. When cleaning is necessary during a journey, this should be carried out with the minimum of stress to the animals.

g) Where an animal handler believes that there is a significant risk of disease among the animals to be loaded or significant doubt as to their fitness to travel, the animals should be examined by a veterinarian.

2. Selection of compatible groups

Compatible groups should be selected before transport to avoid adverse animal welfare consequences. The following guidelines should be applied when assembling groups of animals:

a) animals reared together should be maintained as a group; animals with a strong social bond, such as a dam and offspring, should be transported together;

b) animals of the same species should not be mixed unless there is a significant likelihood of aggression; aggressive individuals should be segregated (recommendations for specific species are described in detail in Article 3.7.3.10.). For some species, animals from different groups should not be mixed because poor welfare occurs unless they have established a social structure;

c) young or small animals should be separated from older or larger animals, with the exception that dam and offspring should be transported together of nursing mothers with young at foot;

d) animals with horns or antlers should not be mixed with animals lacking horns or antlers unless judged to be compatible;

e) animals of different species should not be mixed unless they are judged to be compatible.

3. Shelter in the assembly/holding area

Assembly/holding areas should be designed to:

a) securely hold the animals;

b) maintain a safe environment from hazards, including predators and disease;

c) protect animals from exposure to severe weather conditions;

d) allow for maintenance of social groups, and

e) allow for rest, and appropriate water and feed.

4. Effect of travel experience, long and short term

a) Consideration should be given to an animal’s previous transport experience, training and conditioning as these may reduce fear and stress in animals. Animals that are carefully and regularly transported may show less adverse responses to transport.

b) Exposure to familiar personnel should reduce the fearfulness of animals and improve their approachability during transport procedures.
5. **Fitness to travel**

a) Each animal should be inspected by a veterinarian or an *animal handler* to assess fitness to travel. If its fitness to travel is in doubt, the animal should be examined by a veterinarian. Animals found unfit to travel should not be loaded onto a vehicle, except for transport to receive veterinary treatment.

b) Humane and effective arrangements should be made by the owner or agent for the handling and care of any animal rejected as unfit to travel.

c) Animals that are unfit to travel include:

   i) those that are sick, injured, weak, disabled or fatigued;
   
   ii) those that are unable to stand unaided and bear weight on each leg;
   
   iii) those that are blind in both eyes;
   
   iv) those that cannot be moved without causing them additional suffering;
   
   v) newborn with an unhealed navel;
   
   vi) pregnant animals which are likely to give birth during the journey and those that would be in the final 10% of their gestation period at the planned time of unloading;
   
   vii) females travelling without young which have given birth within the previous 48 hours;
   
   viii) those whose body condition would result in poor welfare because of the expected climatic conditions.

d) Risks during transport can be reduced by selecting animals best suited to the conditions of travel and those that are acclimatised to expected weather conditions.

e) Animals ‘at risk’ which require special conditions (such as in the design of facilities and vehicles, and the length of the journey) and additional attention during transport, may include:

   i) large or obese individuals;
   
   ii) very young or old animals;
   
   iii) excitable or aggressive animals;
   
   iv) animals which have had little contact with humans;
   
   v) animal subject to motion sickness;
   
   vi) females in late pregnancy or heavy lactation, dam and offspring;
   
   vii) *those animals* with a history of exposure to stressors or pathogenic agents prior to transport.
Appendix XXI (contd)

6. Specific species requirements

Transport procedures should be able to take account of variations in the behaviour of the species. Flight zones, social interactions and other behaviour vary significantly among species and even within species. Facilities and handling procedures that are successful with one species are often ineffective or dangerous with another.

Recommendations for specific species are described in detail in Article 3.7.3.10.

Article 3.7.3.6.

Loading

1. Experienced Competent supervision
   a) Since loading has been shown to be the procedure most likely to be the cause of poor welfare in transported animals, the methods to be used should be carefully planned. Loading should be carefully planned as it has the potential to be the cause of poor welfare in transported animals.
   b) Loading should be supervised and/or conducted by animal handlers. These animal handlers should ensure that animals are loaded quietly and without unnecessary noise, harassment or force, and that untrained assistants or spectators do not impede the process.
   c) When containers are loaded onto a vehicle, this should be carried out in such a way to avoid poor animal welfare.

2. Facilities
   a) The facilities for loading including the collecting area, races and loading ramps should be designed and constructed to take into account the needs and abilities of the animals with regard to dimensions, slopes, surfaces, absence of sharp projections, flooring, etc.
   b) Loading facilities should be properly illuminated to allow the animals to be observed by the animal handler(s), and to allow the animals’ ease of movement at all times. Facilities should provide uniform lighting light levels directly over approaches to sorting pens, chutes, loading ramps, with brighter lighting light levels inside vehicles / containers, in order to minimise baulking. Dim lighting light levels may be advantageous for the catching of poultry and some other animals. Artificial lightening may be required.
   c) Ventilation during loading and the journey should provide for fresh air, the removal of excessive heat, humidity and noxious fumes (such as ammonia and carbon monoxide), and the prevention of accumulations of ammonia and carbon dioxide. Under warm and hot conditions, ventilation should allow for the adequate convective cooling of each animal. In some instances, adequate ventilation can be achieved by increasing the space allowance for animals.

3. Goads and other aids

The following principles should apply:
Appendix XXI (contd)

a) Animals which have little or no room to move should not be subjected to physical force or goads and other aids which compel movement.

b) Useful and permitted aids include panels, flags, plastic paddles, flappers (a length of cane with a short strap of leather or canvas attached), plastic bags and metallic rattles; they should be used in a manner sufficient to encourage and direct movement of the animals but without physical contact with them.

c) Painful procedures (including whipping, tail twisting, use of nose twitches, pressure on eyes, ears or external genitalia), or the use of unsuitable goads or other aids (including sticks with sharp ends, lengths of metal piping, fencing wire or heavy leather belts), should not be used to move animals.

d) The use of goads which administer electric shocks should be discouraged, and restricted to that necessary to assist movement of the animal. Such use should be limited to battery-powered goads on the hindquarters of adult pigs and cattle, and never on sensitive areas such as the eyes, mouth, ears, anogenital region or belly. Such instruments should not be used on other animals.

e) The use of well trained dogs to help with the loading of some species may be acceptable.

f) The throwing or dropping of animals, or their lifting or dragging by body parts such as their tail, head, horns, ears, limbs, wool, hair or feathers, should not be permitted. The manual lifting of small animals is permissible.

g) Shouting or yelling at animals or making loud noises e.g. through the cracking of whips to encourage them to move should not occur, as such actions may make the animals agitated, leading to crowding or falling.

Article 3.7.3.7.

Travel

1. General considerations

a) Drivers and animal handlers should check the load immediately before departure to ensure that the animals have been properly loaded. Each load should be checked again early in the trip and adjustments made as appropriate. Periodic checks should be made throughout the trip.

b) Drivers should utilise smooth, defensive driving techniques, without sudden turns or stops, to minimise uncontrolled movements of the animals.

2. Methods of restraining or containing animals

a) Methods of restraining animals should be appropriate to the species and age of animals involved and the training of the individual animal.

b) Recommendations for specific species are described in detail in Article 3.7.3.10.
Appendix XXI (contd)

3. Regulating the environment within vehicles or containers

   a) Animals should be protected against harm from hot or cold conditions during travel. Effective ventilation procedures for maintaining the animals’ environment within vehicles or containers will vary according to whether conditions are cold, hot and dry or hot and humid, but in all conditions a build-up of noxious gases should be prevented. Specific temperature and humidity parameters are described in detail in Appendix XXX.

   b) The animals’ environment in hot weather can be regulated by the flow of air produced by the movement of the vehicle. In warm and hot weather, the duration of journey stops should be minimised and vehicles should be parked under shade, with *maximal adequate and appropriate* ventilation.

   c) To minimise slipping and soiling, and maintain a healthy environment, urine and faeces should be removed from floors when necessary and disposed of in such a way as to prevent the transmission of disease and in compliance with all relevant health and environmental legislation.

4. Sick, injured and dead animals

   a) A driver or *animal handler* finding sick, injured or dead animals should act according to a predetermined emergency response plan.

   b) If possible, sick or injured animals should be segregated.

   c) Ferries (roll-on roll-off) should have procedures to treat sick or injured animals during the journey.

   d) In order to reduce the likelihood that animal transport will increase the spread of infectious disease, contact between transported animals, or the waste products of the transported animals, and other farm animals should be minimised.

   e) During the journey, when disposal of a dead animal becomes necessary, this should be carried out in such a way as to prevent the transmission of disease and in compliance with all relevant health and environmental legislation.

   f) When euthanasia is necessary, the driver or *animal handler* should ensure that it is carried out as *humanely* and results in immediate death. When necessary, assistance should be sought from a veterinarian or other person(s) competent in *humane* euthanasia procedures. Recommendations for specific species are described in Appendix 3.7.6. on humane killing of animals for disease control purposes.

5. Water and feed requirements

   a) If journey duration is such that feeding or watering is required or if the species requires feed or water throughout, access to suitable feed and water for all the animals (*appropriate for their species and age*) carried in the vehicle should be provided. There should be adequate space for all animals to move to the feed and water sources and due account taken of likely competition for feed.

   b) Recommendations for specific species are described in detail in Article 3.7.3.10.
6. **Rest periods and conditions including hygiene**
   
a) Animals that are being transported should be rested at appropriate intervals during the journey and offered feed and water, either on the vehicle or, if necessary, unloaded into suitable facilities.

b) Suitable facilities should be used en route, when resting requires the unloading of the animals. These facilities should meet the needs of the particular animal species and should allow access of all animals to feed and water.

7. **In-transit observations**
   
a) Animals being transported by road should be observed soon after a journey is commenced and whenever the driver has a rest stop (with a maximum interval of 5 hours). After meal breaks and refuelling stops, the animals should be observed immediately prior to departure.

b) Animals being transported by rail should be observed at each scheduled stop nearest to 5 hours since the last observation. The responsible rail transporter should monitor the progress of trains carrying animals and take all appropriate action to minimise delays.

c) During stops, it should be ensured that the animals continue to be properly confined, have appropriate feed and water, and their physical condition is satisfactory.

   **Article 3.7.3.8.**

**Unloading and post-journey handling**

1. **General considerations**
   
a) The required facilities and the principles of animal handling detailed in Article 3.7.3.6. apply equally to unloading, but consideration should be given to the likelihood that the animals will be fatigued.

b) Unloading should be supervised and/or conducted by an animal handler with knowledge and experience of the behavioural and physical characteristics of the species being unloaded. Animals should be unloaded from the vehicle into appropriate facilities as soon as possible after arrival at the destination but sufficient time should be allowed for unloading to proceed quietly and without unnecessary noise, harassment or force.

c) Facilities should provide all animals with appropriate care and comfort, adequate space and ventilation, access to feed (if appropriate) and water, and shelter from extreme weather conditions.

d) For details regarding the unloading of animals at a slaughterhouse, see Appendix 3.7.5. on slaughter of animals for human consumption.

2. **Sick and injured animals**
   
a) An animal that has become sick, injured or disabled during a journey should be appropriately treated or humanely killed (see Appendix 3.7.6. on humane killing of animals for disease control purposes). When necessary, veterinary advice should be sought in the care and treatment of these animals. In some cases, where animals are non-ambulatory due to fatigue, injury or sickness, it may be in the best welfare interests of the animal to be treated or euthanased aboard the vehicle.
Appendix XXI (contd)

b) At the destination, the animal handler during transit should ensure that responsibility for the welfare of sick, injured or disabled animals is transferred to a suitable person.

c) There should be appropriate facilities and equipment for the humane unloading of animals that are non-ambulatory due to fatigue, injury or sickness. These animals should be unloaded in a manner that causes the least amount of suffering. After unloading, separate pens and other appropriate facilities should be available for sick or injured animals.

d) Feed, if appropriate, and water should be available for each sick or injured animal.

3. Addressing disease risks

The following should be taken into account in addressing the greater risk of disease due to animal transport and the possible need for segregation of transported animals at the destination:

a) increased contact among animals, including those from different sources and with different disease histories;

b) increased shedding of pathogens and increased susceptibility to infection related to stress and impaired defences against disease, including immunosuppression;

c) exposure of animals to pathogens which may contaminate vehicles, resting points, markets, etc.

4. Cleaning and disinfection

a) Vehicles, crates, containers, etc. used to carry the animals should be cleaned before re-use through the physical removal of manure and bedding by scraping, washing and flushing vehicles and containers with water and detergent. This should be followed by disinfection when there are concerns about disease transmission.

b) Manure, litter and bedding should be disposed of in such a way as to prevent the transmission of disease and in compliance with all relevant health and environmental legislation.

c) When disposal of a dead animal becomes necessary, this should be carried out in such a way as to prevent the transmission of disease and in compliance with all relevant health and environmental legislation.

b) Manure, litter, bedding and the bodies of any animals which die during the journey should be disposed of in such a way as to prevent the transmission of disease and in compliance with all relevant health and environmental legislation.

c) Establishments like livestock markets, slaughterhouses, resting sites, railway stations, etc. where animals are unloaded should be provided with appropriate areas for the cleaning and disinfection of vehicles.

d) Where disinfection is necessary, it should be carried out with the minimum stress to the animals.

Article 3.7.3.9.

Actions in the event of a refusal to allow the completion of the journey

1. The welfare of the animals should be the first consideration in the event of a refusal to allow the completion of the journey.
2. When the animals have been refused import, the Competent Authority of that country should make available suitable isolation facilities to allow the unloading of animals from a vehicle and their secure holding, without posing a risk to the health of national herd or flock, pending resolution of the situation. In this situation, the priorities should be:

a) the Competent Authority of the importing country should provide urgently in writing the reasons for the refusal;

b) in the event of a refusal for animal health reasons, the Competent Authority of the importing country should provide urgent access to a veterinarian, where possible an OIE veterinarian(s) appointed by the Director General, to assess the animals’ health status with regard to the importing country’s concerns, and the necessary facilities and approvals to expedite the required diagnostic testing;

c) the Competent Authority of the importing country should provide access to allow continued assessment of the health and other aspects of the welfare of the animals;

d) if the matter cannot be promptly resolved, the Competent Authorities of the exporting and importing countries should call on the OIE to mediate.

3. In the event that a Competent Authority requires the animals to remain on the vehicle, the priorities should be:

a) the Competent Authority should allow reprovisionsing of the vehicle with water and feed as necessary;

b) the Competent Authority should provide urgently in writing the reasons for the refusal;

c) in the event of a refusal for animal health reasons, the Competent Authority should provide urgent access to an independent veterinarian(s) to assess the animals’ health status, and the necessary facilities and approvals to expedite the required diagnostic testing;

d) the Competent Authority should provide access to allow continued assessment of the health and other aspects of the welfare of the animals, and the necessary actions to deal with any animal issues which arise.

4. The OIE should utilise its dispute settlement mechanism to identify a mutually agreed solution which will address animal health and any other welfare issues in a timely manner.

Article 3.7.3.10.

Species specific issues

(To be developed)
APPENDIX 3.7.5.

GUIDELINES FOR THE SLAUGHTER OF ANIMALS FOR HUMAN CONSUMPTION

Article 3.7.5.1.

General principles

1. Object

These guidelines address the need to ensure the welfare of food animals during pre-slaughter and slaughter processes, until they are dead.

These guidelines apply to the slaughter in slaughterhouses of the following domestic animals commonly slaughtered in slaughterhouses, that is: cattle, buffalo, sheep, goats, deer, horses, pigs, ratites and poultry. Other animals, wherever they have been reared, and all animals slaughtered outside slaughterhouses should be managed to ensure that their transport, lairaging, restraint and slaughter is carried out without causing undue stress to the animals; the principles underpinning these guidelines apply also to these animals.

2. Personnel

Persons engaged in the unloading, moving, lairaging, care, restraining, stunning, slaughter and bleeding of animals play an important role in the welfare of those animals. For this reason, there should be a sufficient number of personnel, who should be patient, considerate, competent and familiar with the guidelines outlined in the present Appendix and their application within the national context.

Competence may be gained through formal training and/or practical experience. This competence should be demonstrated through a current certificate from an independent body accredited by the Competent Authority.

The management of the slaughterhouse and the Veterinary Services should ensure that slaughterhouse staff are competent and carry out their tasks in accordance with the principles of animal welfare.

The management of the slaughterhouse and the Veterinary Services should ensure that slaughterhouse staff carry out their tasks in accordance with the principles of animal welfare.

3. Animal behaviour

Animal handlers should be experienced and competent in handling and moving farm livestock, and understand the behaviour patterns of animals and the underlying principles necessary to carry out their tasks.
Appendix XXII (contd)

The behaviour of individual animals or groups of animals will vary, depending on their breed, sex, temperament and age and the way in which they have been reared and handled. Despite these differences, the following behaviour patterns which are always present to some degree in domestic animals, should be taken into consideration in handling and moving the animals.

Most domestic livestock are kept in herds and follow a leader by instinct.

Animals which are likely to be hostile to each other in a group situation should not be mixed at slaughterhouses.

The desire of some animals to control their personal space should be taken into account in designing facilities.

Domestic animals will try to escape if an animal handler approaches closer than a certain distance. This critical distance, which defines the flight zone, varies among species and individuals of the same species, and depends upon previous contact with humans. Animals reared in close proximity to humans i.e. tame have a small flight zone, whereas those kept in free range or extensive systems may have flight zones which may vary from one metre to many metres. Animal handlers should avoid sudden penetration of the flight zone which may cause a panic reaction which could lead to aggression or attempted escape.

An example of a flight zone (cattle)
Animal handlers should use the point of balance at an animal’s shoulder to move animals, adopting a position behind the point of balance to move an animal forward and in front of the point of balance to move it backward.

Domestic animals have wide-angle vision but only have limited forward binocular vision and poor perception of depth. This means that they can detect objects and movements beside and behind them, but can only judge distances directly ahead.

Although all domestic animals have a highly sensitive sense of smell, they react in different ways to the smells of slaughterhouses. Smells which cause fear or other negative responses should be taken into consideration when managing animals.

Domestic animals can hear over a greater range of frequencies than humans and are more sensitive to higher frequencies. They tend to be alarmed by constant loud noise and by sudden noises, which may cause them to panic. Sensitivity to such noises should also be taken into account when handling animals.

4. Distractions and their removal

Distractions that may cause approaching animals to stop, baulk or turn back should be designed out from new facilities or removed from existing ones. Below are examples of common distractions and methods for eliminating them:

a) reflections on shiny metal or wet floors - move a lamp or change lighting;

b) dark entrances to chutes, races, stun boxes or conveyor restrainers - illuminate with indirect lighting which does not shine directly into the eyes of approaching animals;

c) animals seeing moving people or equipment up ahead - install solid sides on chutes and races or install shields;
Appendix XXII (contd)

d) chains or other loose objects hanging in chutes or on fences - remove them;

e) uneven floors or a sudden drop in floor levels at the entrance to conveyor restrainers – avoid uneven floor surfaces or install a solid false floor under the restrainer to provide an illusion of a solid and continuous walking surface;

f) sounds of air hissing from pneumatic equipment - install silencers or use hydraulic equipment or vent high pressure to the external environment using flexible hosing;

g) clanging and banging of metal objects - install rubber stops on gates and other devices to reduce metal to metal contact;

h) air currents from fans or air curtains blowing into the face of animals - redirect or reposition equipment.

Article 3.7.5.2.

Moving and handling animals

1. General considerations

Animals should be transported to slaughter in a way that minimises adverse animal health and welfare outcomes, and the transport should be conducted in accordance with the OIE guidelines for the transportation of animals (Chapters 3.7.2 and 3.7.3).

The following principles should apply to unloading animals, moving them into lairage pens, out of the lairage pens and up to the slaughter point:

a) The conditions of the animals should be assessed upon their arrival for any animal welfare and health problems.

b) Injured or sick animals, requiring immediate slaughter, should be killed humanely, preferably at the site where they are found in accordance with the OIE guidelines for the killing of animals for disease control purposes (Chapter 3.7.6).

c) The use of force on animals that have little or no room to move should not occur.

d) The use of instruments which administer electric shocks (e.g. goads and prods) and their power output should be restricted to that necessary to assist movement of all the animals, and only when an animal has a clear path ahead to move. If such use is necessary, it should be limited to the hindquarters of pigs and large ruminants, and never on sensitive areas such as the eyes, mouth, ears, anogenital region or belly. Such instruments should not be used on horses, sheep and goats of any age, or on calves or piglets, nor on animals that have little or no room to move.

e) Performance standards should be established in which numerical scoring is used to evaluate the use of such instruments, and to measure the percentage of animals moved with an electric instrument and the percentage of animals slipping or falling at a point in the slaughterhouse; the slaughterhouse should be investigated for faults in flooring, raceway design, lighting or handling, and these should be rectified to enable free movement of the animals without the need to use such instruments.
Performance standards should be established in which numerical scoring is used to evaluate the use of such instruments and to measure the percentage of animals moved with an electric instrument. In properly designed and constructed facilities with competent animal handlers, it should be possible to move 75% or more of the animals without the use of electric instruments.

f) Useful and permitted aids for moving animals include panels, flags, plastic paddles, flappers (a length of cane with a short strap of leather or canvas attached), plastic bags and metallic rattles; they should be used in a manner sufficient to encourage and direct movement of the animals but without physical contact with them. Aids for moving animals such as panels, flags, plastic paddles, flappers (a length of cane with a short strap of leather or canvas attached), plastic bags and metallic rattles should be used in a manner sufficient to encourage and direct movement of the animals.

g) Shouting or yelling at animals or making loud noises e.g. through the cracking of whips to encourage them to move should not occur as such actions may make the animals agitated, leading to crowding or falling.

h) Implements which cause pain and suffering such as large sticks, sticks with sharp ends, metal piping, fencing wire or heavy leather belts should not be used to move animals.

i) Animals should be grasped or lifted in a manner which avoids pain or suffering and physical damage (e.g. bruising, fractures, dislocations). In the case of quadrupeds, manual lifting by a person should only be used in young animals or small species, and in a manner appropriate to the species; grasping or lifting such animals only by their wool, hair, feet, neck, ears or tails causing pain or suffering should not be permitted, except in an emergency where animal welfare or human safety may otherwise be compromised.

j) Conscious animals should not be thrown or dragged.

k) Animals should not be forced to move at a speed greater than their normal walking pace, in order to minimise injury through falling or slipping. Performance standards should be established where numerical scoring of the prevalence of animals slipping or falling is used to evaluate whether animal moving practices and/or facilities should be improved. In properly designed and constructed facilities with competent animal handlers, it should be possible to move 99% of animals without their falling.

l) Animal handlers should not force an animal to walk over the top of other animals. Animals for slaughter should not be forced to walk over the top of other animals.

m) Under no circumstances should animal handlers resort to violent acts to move animals, such as crushing or breaking animals’ tails, grasping animals’ eyes or pulling them by their ears. Animal handlers should never apply an injurious object or irritant substance to animals and especially not to sensitive areas such as eyes, mouth, ears, anogenital region or belly. The throwing or dropping of animals, or their lifting or dragging by body parts such as their tail, head, horns, ears, limbs, wool, hair or feathers, should not be permitted. The manual lifting of small animals is permissible.

2. Provisions relevant to animals delivered in containers

a) Containers in which animals are transported should be handled with care, and should not be thrown, dropped or knocked over. Where possible, they should be loaded and unloaded horizontally and mechanically.
Appendix XXII (contd)

b) Animals delivered in containers with perforated or flexible bottoms should be unloaded with particular care in order to avoid injury. Where appropriate, animals should be unloaded from the containers individually.

c) Animals which have been transported in containers should be slaughtered as soon as possible; mammals and ratites which are not taken directly upon arrival to the place of slaughter should have drinking water available to them from appropriate facilities at all times. Delivery of poultry for slaughter should be scheduled such that they are not deprived of water at the premises for longer than 12 hours. Animals which have not been slaughtered within 12 hours of their arrival should be fed, and should subsequently be given moderate amounts of food at appropriate intervals.

3. Provisions relevant to restraining and containing animals

a) Provisions relevant to restraining animals for stunning or slaughter without stunning, to help maintain animal welfare, include:

   i) provision of a non-slip floor;

   ii) avoidance of excessive pressure applied by restraining equipment that causes struggling or vocalisation in animals;

   iii) equipment engineered to reduce noise of air hissing and clanging metal;

   iv) absence of sharp edges in restraining equipment that would harm animals;

   v) avoidance of jerking or sudden movement of restraining device.

b) Methods of restraint causing avoidable suffering, such as the following, should not be used in conscious animals because they cause severe pain and stress:

   i) suspending or hoisting animals (other than poultry) by the feet or legs;

   ii) indiscriminate and inappropriate use of stunning equipment;

   iii) mechanical clamping of an animal's legs or feet (other than shackles used in poultry and ostriches) as the sole method of restraint;

   iv) breaking legs, cutting leg tendons or blinding animals in order to immobilise them;

   v) severing the spinal cord, for example using a puntilla or dagger, to immobilise animals using electric currents to immobilise animals, except for proper stunning.

Article 3.7.5.3.

Lairage design and construction

1. General considerations

The lairage should be designed and constructed to hold an appropriate number of animals in relation to the throughput rate of the slaughterhouse without compromising the welfare of the animals.
In order to permit operations to be conducted as smoothly and efficiently as possible without injury or undue stress to the animals, the lairage areas should be designed and constructed so as to allow the animals to move freely in the required direction, using their behavioural characteristics and without undue penetration of their flight zone.

The following guidelines may help to achieve this.

2. **Design of lairages**

a) The lairage should be designed to allow a one-way flow of animals from unloading to the point of slaughter, with a minimum number of abrupt corners to negotiate.

b) In red meat slaughterhouses, pens, passageways and races should be arranged in such a way as to permit inspection of animals at any time, and to permit the removal of sick or injured animals when considered to be appropriate, for which separate appropriate accommodation should be provided.

c) Each animal should have room to stand up and lie down and, when confined in a pen, to turn around. The lairage should have sufficient accommodation for the number of animals intended to be held. Drinking water should always be available to the animals, and the method of delivery should be appropriate to the type of animal held. Troughs should be designed and installed in such a way as to minimise the risk of fouling by faeces, without introducing risk of bruising and injury in animals, and should not hinder the movement of animals.

d) Holding pens should be designed rectangular rather than square, to allow as many animals as possible to stand or lie down against a wall. Where feed troughs are provided, they should be sufficient in number and feeding space to allow adequate access of all animals to feed. The feed trough should not hinder the movement of animals.

e) Where tethers, ties or individual stalls are used, these should be designed so as not to cause injury or distress, especially when the animals are lying down, standing up, drinking and feeding to the animals and should also allow the animals to stand, lie down and access any food or water that may need to be provided.

f) Passageways and races should be either straight or slightly consistently curved, as appropriate to the animal species. Passageways and races should have solid sides, but when there is a double race, the shared partition should allow adjacent animals to see each other. For pigs and sheep, passageways should be wide enough to enable two or more animals to walk side by side for as long as possible. At the point where passageways are reduced in width, this should be done by a means which prevents excessive bunching of the animals.

g) Animal handlers should be positioned alongside races and passageways on the inside radius of any curve, to take advantage of the natural tendency of animals to circle an intruder. Where one-way gates are used, they should be of a design which avoids bruising. Races should be horizontal but where there is a slope, they should be constructed to allow the free movement of animals without injury.

h) There should be a waiting pen, with a level floor and solid sides, between the holding pens and the race leading to the point of stunning or slaughter, to ensure a steady supply of animals for stunning or slaughter and to avoid having animal handlers trying to rush animals from the holding pens. The waiting pen should preferably be circular, but in any case, so designed that animals cannot be trapped or trampled.
Appendix XXII (contd)

i) Ramps or lifts should be used for loading and unloading of animals where there is a difference in height or a gap between the floor of the vehicle and the unloading area. Unloading ramps should be designed and constructed so as to permit animals to be unloaded from vehicles on the level or at the minimum gradient achievable. Lateral side protection should be available to prevent animals escaping or falling. The ramp should be well drained, non-slippery, with secure footholds, and adjustable to facilitate easy movement of animals without causing distress or injury.

3. Construction of lairages

a) Lairages should be constructed and maintained so as to provide protection from unfavourable climatic conditions, using strong and resistant materials such as concrete and metal which has been treated to prevent corrosion. Surfaces should be easy to clean. There should be no sharp edges or protuberances which may injure the animals.

b) Floors should be well drained and not slippery; they should not cause injury to the animals' feet. Where necessary, floors should be insulated or provided with appropriate bedding. Drainage grids should be placed at the sides of pens and passageways and not where animals would have to cross them. Discontinuities or changes in floor patterns or texture which could cause baulking in the movement of animals should be avoided.

c) Lairages should be provided with adequate lighting, but care should be taken to avoid harsh lights and shadows, which frighten the animals or affect their movement. The fact that animals will move more readily from a darker area into a well-lit area might be exploited by providing for lighting that can be regulated accordingly.

d) Lairages should be well ventilated, and the air flow should be arranged so that odours and draughts do not adversely affect the health and welfare of the animals adequately ventilated to ensure that waste gases, e.g. ammonia do not build up and that draughts at animal height are minimised. Ventilation should be able to cope with the range of expected climatic conditions and the number of animals the lairage will be expected to hold.

e) Care should be taken to protect the animals from excessively or potentially disturbing noises, for example by avoiding the use of noisy hydraulic or pneumatic equipment, and muffling noisy metal equipment by the use of suitable padding, or by minimising the transmission of such noise to the areas where animals are held and slaughtered.

f) Where animals are kept in outdoor lairages without natural shelter or shade, they should be protected from the effects of adverse weather conditions.

Article 3.7.5.4.

Care of animals in lairages

Animals in lairages should be cared for in accordance with the following guidelines:

1. As far as possible, established groups of animals should be kept together. Each animal should have enough space to stand up, lie down and turn around. Animals hostile to each other should be separated.
2. Where tethers, ties or individual stalls are used, they should allow animals to stand up and lie down without causing injury or distress.

3. Where bedding is provided, it should be maintained in a condition that minimises risks to the health and safety of the animals, and sufficient bedding should be used so that animals do not become soiled with manure.

4. Animals should be kept securely in the lairage, and care should be taken to prevent them from escaping and from predators.

5. Suitable drinking water should be available to the animals on their arrival and at all times to animals in lairages unless they are to be slaughtered without delay.

6. If animals are not to be slaughtered as soon as possible, suitable feed should be available to the animals on arrival and at intervals appropriate to the species. Unweaned animals should be slaughtered as soon as possible.

7. In order to prevent heat stress, animals subjected to high temperatures, particularly pigs and poultry, should be cooled by the use of water sprays, fans or other suitable means. However, the potential for water sprays to reduce the ability of animals to thermoregulate (especially poultry) should be considered in any decision to use water sprays.

8. The lairage area should be well lit in order to enable the animals to see clearly without being dazzled. During the night, the lights should be dimmed. Lighting should also be adequate to permit inspection of all animals. Subdued lighting, and for example, blue light may be useful in poultry lairages in helping to calm birds.

9. The condition and state of health of the animals in a lairage should be inspected at least every morning and evening by a veterinarian or, under the latter's responsibility, by another competent person. Animals which are sick, weak, injured or showing visible signs of distress should be treated or humanely killed immediately.

10. Lactating dairy animals should be slaughtered as soon as possible. Dairy animals with obvious udder distension should be milked to minimise udder discomfort.

11. Pregnant animals giving birth during the journey or in the lairage should be slaughtered as soon as possible or provided with conditions which are appropriate for suckling for its welfare and the welfare of the newborn. Under normal circumstances, animals which are expected to give birth during a journey should not be transported.

12. Animals with horns or tusks capable of injuring other animals, if aggressive, should be penned separately.

Recommendations for specific species are described in detail in Articles 3.7.5.5. to 3.7.5.8.
Management of foetuses during slaughter of pregnant animals

The welfare of foetuses during slaughter of pregnant animals needs to be safeguarded.

1. Foetuses should not be removed from the uterus sooner than five minutes after the maternal neck or chest cut, to ensure absence of consciousness. A foetal heartbeat will usually still be present and foetal movements may occur at this stage, but these are only a cause for concern if the exposed foetus successfully breathes air.

2. If a live mature foetus is removed from the uterus, it should be prevented from inflating its lungs and breathing air (e.g. by clamping the trachea).

3. When uterine, placental or foetal tissues, including foetal blood, are not to be collected as part of the post-slaughter processing of pregnant animals, all foetuses should be left inside the unopened uterus until they are dead. When uterine, placental or foetal tissues are to be collected, where practical, foetuses should not be removed from the uterus until at least 15-20 minutes after the maternal neck or chest cut.

4. If there is any doubt about consciousness, the foetus should be killed with a captive bolt or a blow to the head with a suitable blunt instrument.

The above guidelines do not refer to foetal rescue. Foetal rescue, the practice of attempting to revive foetuses found alive at evisceration of the dam, should not be attempted during normal commercial slaughter as it may lead to serious welfare complications in the newborn animal. These include impaired brain function resulting from oxygen shortage before rescue is completed, compromised breathing and body heat production because of foetal immaturity, and an increased incidence of infections due to a lack of colostrum.
### Article 3.7.5.6.

#### Summary of acceptable handling and restraining methods and the associated animal welfare issues

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<td>Poultry</td>
<td>Proper design and operation of equipment</td>
<td>Stress and injury due to tipping in dump-module systems; height of tipping conscious poultry broken bones and dislocations</td>
<td>Presentation of birds for shackling prior to electrical stunning Gas stunning</td>
<td>Flat bed/deck Tipped out of containers on to conveyors</td>
<td>Mechanical - upright</td>
<td></td>
</tr>
<tr>
<td>Poultry</td>
<td>Competent animal handlers; proper design and operation of equipment</td>
<td>Inversion stress; pain from compression on leg bones</td>
<td>Electrical stunning Slaughter without stunning</td>
<td>Poultry shackle</td>
<td>Suspension and/or inversion</td>
<td></td>
</tr>
<tr>
<td>Poultry</td>
<td>Competent animal handlers; proper design and operation of equipment</td>
<td>Inversion stress</td>
<td>Electrical – head-only Captive bolt Slaughter without stunning</td>
<td>Cone</td>
<td>Suspension and/or inversion</td>
<td></td>
</tr>
<tr>
<td>Ostriches</td>
<td>Competent animal handlers; proper equipment design and operation</td>
<td>Stress of resisting restraint in ostriches</td>
<td>Electrical – head-only</td>
<td>Mechanical leg clamping</td>
<td>Upright restraint</td>
<td></td>
</tr>
</tbody>
</table>
### Summary of acceptable handling and restraining methods and the associated animal welfare issues (contd)

<table>
<thead>
<tr>
<th>Presentation of animals</th>
<th>Specific procedure</th>
<th>Specific purpose</th>
<th>AW concerns/implications</th>
<th>Key AW requirements</th>
<th>Applicable species</th>
</tr>
</thead>
<tbody>
<tr>
<td>Restraining by inversion</td>
<td>Rotating box</td>
<td>Fixed side(s) (e.g. Weinberg pen)</td>
<td>Slaughter without stunning</td>
<td>Inversion stress; stress of resisting restraint, prolonged restraint, inhalation of blood and ingesta. Keep restraint as brief as possible</td>
<td>Proper design and operation of equipment</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Compressible side(s)</td>
<td>Slaughter without stunning</td>
<td>Inversion stress, stress of resisting restraint, prolonged restraint Preferable to rotating box with fixed sides Keep restraint as brief as possible</td>
<td>Proper design and operation of equipment</td>
</tr>
<tr>
<td>Body restraint</td>
<td>Casting/ hobbling</td>
<td>Manual</td>
<td>Mechanical stunning methods Slaughter without stunning</td>
<td>Stress of resisting restraint; animal temperament; bruising Keep restraint as short as possible</td>
<td>Competent animal handlers</td>
</tr>
<tr>
<td>Leg restraints</td>
<td>Rope casting</td>
<td>Mechanical stunning methods Slaughter without stunning</td>
<td>Slaughter without stunning</td>
<td>Stress of resisting restraint; prolonged restraint, animal temperament; bruising Keep restraint as short as possible</td>
<td>Competent animal handlers</td>
</tr>
<tr>
<td></td>
<td>Tying of 3 or 4 legs</td>
<td>Mechanical stunning methods Slaughter without stunning</td>
<td>Slaughter without stunning</td>
<td>Stress of resisting restraint; prolonged restraint, animal temperament; bruising Keep restraint as short as possible</td>
<td>Competent animal handlers</td>
</tr>
</tbody>
</table>
Appendix XXII (contd)

Article 3.7.5.7.

Stunning methods

1. General considerations

The competence of the operators, and the appropriateness, and effectiveness of the method used for stunning and the maintenance of the equipment are the responsibility of the management of the slaughterhouse, and should be checked regularly by a Competent Authority.

Persons carrying out stunning should be properly trained and competent, and should ensure that:

a) the animal is adequately restrained;

b) animals in restraint are stunned as soon as possible;

c) the equipment used for stunning is maintained and operated properly in accordance with the manufacturer's recommendations, in particular with regard to the species and size of the animal;

d) the instrument is applied correctly;

e) stunned animals are bled out (slaughtered) as soon as possible;

f) animals should not be stunned when slaughter is likely to be delayed;

g) backup stunning devices are available for immediate use if the primary method of stunning fails.

In addition, such persons should be able to recognise when an animal is not correctly stunned and should take appropriate action.

2. Mechanical stunning

A mechanical device should be applied usually to the front of the head and perpendicular to the bone surface. The following diagrams illustrate the proper application of the device for certain species.

Cattle

The optimum position for cattle is at the intersection of two imaginary lines drawn from the rear of the eyes to the opposite horn buds.
Appendix XXII (contd)

**Pigs**

The optimum position for pigs is on the midline just above the eyes level, with and directing the shot down the line of the spinal cord.

**Sheep**

The optimum position for hornless sheep and goats is on the midline just above the eye level, and directing the shot down the line of the spinal cord.

**Goats**

The optimum position for heavily horned sheep and horned goats is behind the poll, aiming towards the angle of the jaw.
Place the muzzle. The optimum position for horses is at right angles to the frontal surface, well above the point where imaginary lines from eyes to ears cross.

Signs of correct stunning using a mechanical instrument are as follows:

a) the animal collapses immediately and does not attempt to stand up;

b) the body and muscles of the animal become tonic (rigid) immediately after the shot;

c) normal rhythmic breathing stops; and

d) the eyelid is open with the eyeball facing straight ahead and is not rotated.

3. Electrical stunning

a) General considerations

An electrical device should be applied to the animal in accordance with the following guidelines.

Electrodes should be designed, constructed, maintained and cleaned regularly to ensure that the flow of current is optimal and in accordance with manufacturing specifications. They should be placed so that they span the brain. The application of electrical currents which bypass the brain is unacceptable unless the animal has been stunned. The use of a single current leg-to-leg is unacceptable as a stunning method.

If, in addition, it is intended to cause cardiac arrest, the electrodes should either span the brain and immediately thereafter the heart, on the condition that it has been ascertained that the animal is adequately stunned, or span brain and heart simultaneously.

Electrical stunning equipment should not be applied on animals as a means of guidance, movement, restraint or immobilisation, and shall not deliver any shock to the animal before the actual stunning or killing.

Electrical stunning apparatus should be tested prior to application on animals using appropriate resistors or dummy loads to ensure the power output is adequate to stun animals.
Appendix XXII (contd)

The apparatus should incorporate a device which monitors and displays stunning current delivered to the animals.

Appropriate measures, such as removing excess wool or wetting the skin only at the point of contact, can be taken to minimise impedance of the skin and facilitate effective stunning.

The stunning apparatus required for electrical stunning should be provided with adequate power to achieve continuously the minimum current level recommended for stunning as indicate in the table below:

<table>
<thead>
<tr>
<th>Species</th>
<th>Minimum current levels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cattle</td>
<td>1.5 amps</td>
</tr>
<tr>
<td>Calves</td>
<td>1.0 amps</td>
</tr>
<tr>
<td>Pigs</td>
<td>1.25 amps</td>
</tr>
<tr>
<td>Sheep and goats</td>
<td>1.0 amps</td>
</tr>
<tr>
<td>Lambs</td>
<td>0.7 amps</td>
</tr>
<tr>
<td>Ostriches</td>
<td>0.4 amps</td>
</tr>
</tbody>
</table>

In all cases, the correct current level shall be attained within one second of the initiation of stun and maintained at least for between one and three seconds and in accordance with the manufacturer's instructions.

b) Electrical stunning of birds using a waterbath

In the case of birds suspended on a moving line, measures should be taken to ensure that the birds are not wing flapping at the entrance of the stunner. The birds should be secure in their shackle, but there should not be undue pressure on their shanks.

Waterbaths for poultry should be adequate in size and depth for the type of bird being slaughtered, and their height should be adjustable to allow for the head of each bird to be immersed. The electrode immersed in the bath should extend the full length of the waterbath. Birds should be immersed in the bath up to the base of their wings.

The waterbath should be designed and maintained in such a way that when the shackles pass over the water, they are in continuous contact with the earthed rubbing bar.

The control box for the waterbath stunner should incorporate an ammeter which displays the total current flowing through the birds.

The shackle-to-leg contact should be wetted preferably before the birds are inserted in the shackles. In order to improve electrical conductivity of the water it is recommended that salt be added in the waterbath as necessary. Additional salt should be added regularly as a solution to maintain suitable constant concentrations in the waterbath.

Using waterbaths, birds are stunned in groups and different birds will have different impedances. The voltage should be adjusted so that the total current is the required current per bird as shown in the table hereafter, multiplied by the number of birds in the waterbath at the same time. The following values have been found to be satisfactory when employing a 50 Hertz sinusoidal alternating current.
Birds should receive the current for at least 4 seconds.

<table>
<thead>
<tr>
<th>Species</th>
<th>Current (milliamperes per bird)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Broilers</td>
<td>120</td>
</tr>
<tr>
<td>Layers (spent hens)</td>
<td>120</td>
</tr>
<tr>
<td>Turkeys</td>
<td>150</td>
</tr>
<tr>
<td>Ducks and Geese</td>
<td>130</td>
</tr>
</tbody>
</table>

While a lower current may also be satisfactory, the current shall in any case be such as to ensure that unconsciousness occurs immediately and lasts until the bird has been killed by cardiac arrest or by bleeding. When higher electrical frequencies are used, higher currents may be required.

Every effort shall be made to ensure that no conscious or live birds enter the scalding tank.

In the case of automatic systems, until fail-safe systems of stunning and bleeding have been introduced, a manual back-up system should be in place to ensure that any birds which have missed the waterbath stunner and/or the automatic neck-cutter are immediately stunned and/or killed immediately, and they are dead before entering scald tank.

To lessen the number of unstunned birds, reaching neck cutters, steps should be taken to ensure that small birds do not go on the line amongst bigger birds and that these small birds are stunned separately.

4. **Gas stunning (under study)**

   a) Stunning of pigs by exposure to carbon dioxide (CO$_2$)

   The concentration of CO$_2$ for stunning should be preferably 90% by volume but in any case no less than 80% by volume. After entering the stunning chamber, the animals should be conveyed to the point of maximum concentration of the gas as rapidly as possible and be kept until they are dead or brought into a state of insensibility which lasts until death occur due to bleeding. Ideally, pigs should be exposed to this concentration of CO$_2$ for 3 minutes. Sticking should occur as soon as possible after exit from the gas chamber.

   In any case, the concentration of the gas should be such that it minimises as far as possible all stress of the animal prior to loss of consciousness.

   The chamber in which animals are exposed to CO$_2$ and the equipment used for conveying them through it shall be designed, constructed and maintained in such a way as to avoid injury or unnecessary stress to the animals. The animal density within the chamber should be such to avoid stacking animals on top of each others.

   The conveyor and the chamber shall be adequately lit to allow the animals to see their surroundings and, if possible, each other.

   It should be possible to inspect the CO$_2$ chamber whilst it is in use, and to have access to the animals in emergency cases.
The chamber shall be equipped to continuously measure and display register at the point of stunning the CO$_2$ concentration and the time of exposure, and to give a clearly visible and audible warning if the concentration of CO$_2$ falls below the required level.

b) Inert gas mixtures for stunning pigs

Inhalation of high concentrations of carbon dioxide is aversive and can be distressing to animals. Therefore, the use of non-aversive gas mixtures is being developed.

Such gas mixtures include:

i) a maximum of 2% by volume of oxygen in argon, nitrogen or other inert gases, or

ii) to a maximum of 30% by volume of carbon dioxide and a maximum of 2% by volume of oxygen in mixtures with carbon dioxide and argon, nitrogen or other inert gases.

Exposure time to the gas mixtures should be sufficient to ensure that no pigs regain consciousness before death supervenes through bleeding or cardiac arrest is induced.

c) Gas stunning of poultry

The main objective of gas stunning is to avoid the pain and suffering associated with shackling conscious poultry under water bath stunning and killing systems. Therefore, gas stunning should be limited to birds contained in crates or on conveyors only. The gas mixture should be non-aversive to poultry.

Gas stunning of poultry in their transport containers will eliminate the need for live bird handling at the processing plant and all the problems associated with the electrical stunning. Gas stunning of poultry on a conveyor eliminates the problems associated with the electrical water bath stunning.

Live poultry should be conveyed into the gas mixtures either in transport crates or on conveyor belts.

i) Gas mixtures used for stunning poultry include:

- minimum of 2 minutes exposure to 40% carbon dioxide, 30% oxygen and 30% nitrogen, followed by a minimum of one minute exposure to 80% carbon dioxide in air; or

- minimum of 2 minutes exposure to any mixture of argon, nitrogen or other inert gases with atmospheric air and carbon dioxide, provided that the carbon dioxide concentration does not exceed 30% by volume and the residual oxygen concentration does not exceed 2% by volume; or

- minimum of 2 minutes exposure to argon, nitrogen, other inert gases or any mixture of these gases in atmospheric air with a maximum of 2% residual oxygen by volume; or

- minimum of 2 minutes exposure to a minimum of 55% carbon dioxide in air.
Appendix XXII (contd)

ii) Requirements for effective use are as follows:

- compressed gases should be vaporised prior to administration into the chamber and should be at room temperature to prevent any thermal shock. Under no circumstances, should solid gases with freezing temperatures enter the chamber;

- gas mixtures should be humidified;

- appropriate gas concentrations should be monitored and displayed continuously at the level of the birds inside the chamber.

Under no circumstances, should birds exposed to gas mixtures be allowed to regain consciousness. If necessary, the exposure time should be extended.

5. **Bleeding**

From the point of view of animal welfare, animals which are stunned with a reversible method should be bled without delay and in any case within the following time limits:

<table>
<thead>
<tr>
<th>Stunning method</th>
<th>Maximum delay for bleeding to be started</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrical methods and non penetrating captive bolt</td>
<td>20 seconds</td>
</tr>
<tr>
<td>CO₂</td>
<td>60 seconds (after leaving the chamber)</td>
</tr>
</tbody>
</table>

All animals should be bled by incising both carotid arteries, or the vessels from which they arise (e.g. chest stick). However, when the stunning method used causes cardiac arrest, the incision of all of these vessels is not necessary from the point of animal welfare.

It should be possible for staff to observe, inspect and access the animals throughout the bleeding period. Any animal showing signs of recovering consciousness should be restunned.

After incision of the blood vessels, no scalding carcass treatment or dressing procedures should be performed on the animals for at least 30 seconds, or in any case until all brain-stem reflexes have ceased.
### Appendix XXII (contd)

**Article 3.7.5.8.**

**Summary of acceptable stunning methods and the associated animal welfare issues**

<table>
<thead>
<tr>
<th>Method</th>
<th>Specific method</th>
<th>AW concerns/implications</th>
<th>Key AW requirements applicable</th>
<th>Species</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical</td>
<td>Free bullet</td>
<td>Inaccurate targeting and inappropriate ballistics</td>
<td>Accuracy; head shots only correct ballistics; Operator competence; achieving outright kill with first shot</td>
<td>Cattle, calves, buffalo, deer, horses, pigs (boars and sows)</td>
<td>Personnel safety</td>
</tr>
<tr>
<td>Captive bolt -</td>
<td>Penetrating</td>
<td>Inaccurate targeting, velocity and diameter of bolt</td>
<td>Competent operation and maintenance of equipment; restraint; accuracy</td>
<td>Cattle, calves, buffalo, sheep, goats, deer, horses, pigs, camelids, ratites</td>
<td>(Unsuitable for specimen collection from TSE suspects). A back-up gun should be available in the event of an ineffective shot</td>
</tr>
<tr>
<td>Captive bolt -</td>
<td>Non-penetrating</td>
<td>Inaccurate targeting, velocity of bolt, potentially higher failure rate than penetrating captive bolt</td>
<td>Competent operation and maintenance of equipment; restraint; accuracy</td>
<td>Cattle, calves, sheep, goats, deer, pigs, camelids, ratites</td>
<td>Presently available devices are not recommended for young bulls and animals with thick skull</td>
</tr>
<tr>
<td>Manual percussive</td>
<td>Blow</td>
<td>Inaccurate targeting; insufficient power; size of instrument</td>
<td>Competent animal handlers; restraint; accuracy. Not recommended for general use</td>
<td>Young and small mammals, ostriches and poultry</td>
<td>Mechanical devices potentially more reliable. Where manual percussive blow is used, unconsciousness should be achieved with single sharp blow delivered to central skull bones</td>
</tr>
<tr>
<td>Electrical</td>
<td>Split application: 1. across head then head to chest; 2. across head then across chest</td>
<td>Accidental pre-stun electric shocks; electrode positioning; application of a current to the body while animal conscious; inadequate current and voltage</td>
<td>Competent operation and maintenance of equipment; restraint; accuracy</td>
<td>Cattle, calves, sheep, goats and pigs, ratites and poultry</td>
<td>Systems involving repeated application of head-only or head-to-leg with short current durations (&lt;1 second) in the first application should not be used. Where cardiac arrest occurs, the carcass may not be suitable for Halal</td>
</tr>
</tbody>
</table>
## Summary of acceptable stunning methods and the associated animal welfare issues

<table>
<thead>
<tr>
<th>Method</th>
<th>Specific method</th>
<th>AW concerns/implications</th>
<th>Key AW requirements applicable</th>
<th>Species</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrical</td>
<td>Single application: 1. head only; 2. head to body; 3. head to leg</td>
<td>Accidental pre-stun electric shocks; inadequate current and voltage; wrong electrode positioning; recovery of consciousness</td>
<td>Competent operation and maintenance of equipment; restraint; accuracy</td>
<td>Cattle, calves, sheep, goats, pigs, ratites, poultry</td>
<td>Where cardiac arrest occurs, the carcass may not be suitable for Halal</td>
</tr>
<tr>
<td>Waterbath</td>
<td></td>
<td>Restraint, accidental pre-stun electric shocks; inadequate current and voltage; recovery of consciousness</td>
<td>Competent operation and maintenance of equipment</td>
<td>Poultry only</td>
<td>Where cardiac arrest occurs, the carcass may not be suitable for Halal</td>
</tr>
<tr>
<td>Gaseous</td>
<td>CO₂ air/O₂ mixture; CO₂ inert gas mixture</td>
<td>Aversiveness of high CO₂ concentrations, respiratory distress; inadequate exposure</td>
<td>Concentration; duration of exposure; design, maintenance and operation of equipment; stocking density management</td>
<td>Pigs, poultry</td>
<td>Gaseous methods may not be suitable for Halal</td>
</tr>
<tr>
<td>Inert gases</td>
<td></td>
<td>Recovery of consciousness</td>
<td>Concentration; duration of exposure; design, maintenance and operation of equipment; stocking density management</td>
<td>Pigs, poultry</td>
<td>Gaseous methods may not be suitable for Halal</td>
</tr>
</tbody>
</table>
**Article 3.7.5.9.**

**Summary of acceptable slaughter methods and the associated animal welfare issues**

<table>
<thead>
<tr>
<th>Slaughter methods</th>
<th>Specific method</th>
<th>AW concerns / implications</th>
<th>Key requirements</th>
<th>Species</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bleeding out by severance of blood vessels in the neck without stunning</td>
<td>Full frontal cutting across the throat</td>
<td>Failure to cut both common carotid arteries; occlusion of cut arteries.</td>
<td>A very sharp blade or knife, of sufficient length so that the point of the knife remains outside the incision during the cut; the point of the knife should not be used to make the incision. An incision which does not close over the knife during the throat cut.</td>
<td>Cattle, buffalo, horses, camels, sheep, goats, poultry, ratites</td>
<td>This method is applicable to Halal and Kosher slaughter for relevant species.</td>
</tr>
<tr>
<td>Bleeding with prior stunning</td>
<td>Full frontal cutting across the throat</td>
<td>Failure to cut both common carotid arteries; occlusion of cut arteries; pain during and after the cut.</td>
<td>A very sharp blade or knife, of sufficient length so that the point of the knife remains outside the incision during the cut; the point of the knife should not be used to make the incision. An incision which does not close over the knife during the throat cut.</td>
<td>Cattle, buffalo, horses, camels, sheep, goats.</td>
<td></td>
</tr>
<tr>
<td>Neck stab followed by forward cut</td>
<td>Ineffective stunning; failure to cut both common carotid arteries; impaired blood flow; delay in cutting after reversible stunning</td>
<td>Prompt and accurate cutting</td>
<td>Camelids, sheep, goats, poultry, ratites.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neck stab alone</td>
<td>Ineffective stunning; failure to cut both common carotid arteries; impaired blood flow; delay in cutting after reversible stunning</td>
<td>Prompt and accurate cutting</td>
<td>Camelids, sheep, goats, poultry, ratites.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Summary of acceptable slaughter methods and the associated animal welfare issues (contd)

<table>
<thead>
<tr>
<th>Slaughter methods</th>
<th>Specific method</th>
<th>AW concerns / implications</th>
<th>Key requirements</th>
<th>Species</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bleeding with prior stunning (contd)</td>
<td>Chest stick into major arteries or hollow-tube knife into heart</td>
<td>Ineffective stunning; inadequate size of stick wound; inadequate length of sticking knife; delay in sticking after reversible stunning</td>
<td>Prompt and accurate sticking</td>
<td>Cattle, sheep, goats, pigs</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Chest stick into major arteries or hollow-tube knife into heart</td>
<td>Ineffective stunning; inadequate size of stick wound; inadequate length of sticking knife; delay in sticking after reversible stunning</td>
<td>Prompt and accurate sticking</td>
<td>Cattle, sheep, goats, pigs</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Neck skin cut followed by severance of vessels in the neck</td>
<td>Ineffective stunning; inadequate size of stick wound; inadequate length of sticking knife; delay in sticking after reversible stunning</td>
<td>Prompt and accurate cutting of vessels</td>
<td>Cattle</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Automated mechanical cutting</td>
<td>Ineffective stunning; failure to cut and misplaced cuts. Recovery of consciousness following reversible stunning systems</td>
<td>Design, maintenance and operation of equipment; accuracy of cut; manual back-up</td>
<td>Poultry only</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Manual neck cut on one side</td>
<td>Ineffective stunning; recovery of consciousness following reversible stunning systems</td>
<td>Prior non-reversible stunning</td>
<td>Poultry only</td>
<td>N.B. slow induction of unconsciousness under slaughter without stunning</td>
</tr>
<tr>
<td></td>
<td>Oral cut</td>
<td>Ineffective stunning; recovery of consciousness following reversible stunning systems</td>
<td>Prior non-reversible stunning</td>
<td>Poultry only</td>
<td>N.B. slow induction of unconsciousness in non-stun systems</td>
</tr>
</tbody>
</table>
### Appendix XXII (contd)

<table>
<thead>
<tr>
<th>Slaughter methods</th>
<th>Specific method</th>
<th>AW concerns / implications</th>
<th>Key requirements</th>
<th>Species</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bleeding with prior stunning (contd)</td>
<td>Oral cut</td>
<td>Ineffective stunning; recovery of consciousness following reversible stunning systems</td>
<td>Prior non-reversible stunning</td>
<td>Poultry only</td>
<td>N.B. slow induction of unconsciousness in non-stun systems</td>
</tr>
<tr>
<td>Other methods without stunning</td>
<td>Decapitation with a sharp knife</td>
<td>Pain due to loss of consciousness not being immediate</td>
<td></td>
<td>Sheep, goats, poultry</td>
<td>This method is only applicable to Jhatka slaughter</td>
</tr>
<tr>
<td></td>
<td>Manual neck dislocation and decapitation</td>
<td>Pain due to loss of consciousness not being immediate; difficult to achieve in large birds</td>
<td>Neck dislocation should be performed in one stretch to sever the spinal cord</td>
<td>Poultry only</td>
<td>Slaughter by neck dislocation should be performed in one stretch to sever the spinal cord</td>
</tr>
<tr>
<td>Cardiac arrest in a waterbath electric stunner</td>
<td>Bleeding by evisceration</td>
<td></td>
<td>Induction of cardiac arrest</td>
<td>Quail</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bleeding by neck cutting</td>
<td></td>
<td></td>
<td>Poultry</td>
<td></td>
</tr>
</tbody>
</table>

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**Article 3.7.5.10.**

**Methods, procedures or practices unacceptable on animal welfare grounds**

1. The restraining methods which work through immobilisation by injury such as 'puntilla', breaking legs and ‘leg tendon cutting', cause severe pain and stress in animals. Those methods are not acceptable in any species.

2. The use of the electrical stunning method with a single application leg to leg is ineffective and unacceptable in any species, as it is likely to be painful. The animal welfare concerns are:
   a) accidental pre-stun electric shocks;
   b) inadequate current and voltage;
   c) wrong electrode positioning;
   d) recovery of consciousness.

3. The slaughter method of brain stem severance by piercing through the eye socket or skull bone without prior stunning, is not acceptable in any species.

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OIE Terrestrial Animal Health Standards Commission/September 2005
Appendix XXIII

APPENDIX 3.7.6.

GUIDELINES FOR THE KILLING OF ANIMALS FOR DISEASE CONTROL PURPOSES

Article 3.7.6.1.

General principles

These guidelines are based on the premise that a decision to kill the animals has been made, and address the need to ensure the welfare of the animals until they are dead.

1. All personnel involved in the humane killing of animals should have the relevant skills and competencies. Competence may be gained through formal training and/or practical experience. This competence should be demonstrated through a current certificate from an independent body accredited by a Competent Authority.

2. As necessary, operational procedures should be adapted to the specific circumstances operating on the premises and should address, apart from animal welfare, operator safety, biosecurity and environmental aspects.

3. Following the decision to kill the animals, killing should be carried out as quickly as possible and normal husbandry should be maintained until the animals are killed.

4. The handling and movement of animals should be minimised and when done, it should be done in accordance with the guidelines described below.

5. Animal restraint should be sufficient to facilitate effective killing, and in accordance with animal welfare and operator safety requirements; when restraint is required, killing should follow with minimal delay.

6. When animals are killed for disease control purposes, methods used should result in immediate death or immediate loss of consciousness lasting until death; when loss of consciousness is not immediate, induction of unconsciousness should be non-aversive and should not cause anxiety, pain, distress or suffering in the animals.

7. For animal welfare considerations, young animals should be killed before older animals; for biosecurity considerations, infected animals should be killed first, followed by in-contact animals, and then the remaining animals.

8. There should be continuous monitoring of the procedures by the Competent Authorities to ensure they are consistently effective with regard to animal welfare, operator safety and biosecurity.

9. When the operational procedures are concluded, there should be a written report describing the practices adopted and their effect on animal welfare, operator safety and biosecurity.
Appendix XXIII (contd)

10. To the extent possible to minimise public distress, killing of animals and carcass disposal should be carried out away from public view.

11. These general principles should also apply when animals need to be killed for other purposes such as after natural disasters.

Article 3.7.6.2.

Organisational structure

Disease control contingency plans should be in place at a national level and should contain details of management structure, disease control strategies and operational procedures; animal welfare considerations should be addressed within these disease control contingency plans. The plans should also include a strategy to ensure that an adequate number of personnel trained competent in the humane killing of animals is available. Local level plans should be based on national plans and be informed by local knowledge.

Disease control contingency plans should address the animal welfare issues that may result from animal movement controls.

The operational activities should be led by an official veterinarian who has the authority to appoint the personnel in the specialist teams and ensure that they adhere to the required animal welfare and biosecurity standards. When appointing the personnel, he/she should ensure that the personnel involved has the required competencies.

The official veterinarian should be responsible for all activities across one or more affected premises and should be supported by coordinators for planning (including communications), operations and logistics to facilitate efficient operations.

The official veterinarian should provide overall guidance to personnel and logistic support for operations on all affected premises to ensure consistency in adherence to the OIE animal welfare and animal health guidelines.

A specialist team, led by a team leader answerable to the official veterinarian, should be deployed to work on each affected premises. The team should consist of personnel with the competencies to conduct all required operations; in some situations, personnel may be required to fulfil more than one function. Each team should contain a veterinarian or have access to veterinary advice at all times.

In considering the animal welfare issues associated with killing animals, the key personnel, their responsibilities and competencies required are described in Article 3.7.6.3.

Article 3.7.6.3.

Responsibilities and competencies of the specialist team

1. Team leader

a) Responsibilities

i) plan overall operations on an affected premises;
Appendix XXIII (contd)

ii) determine and address requirements for animal welfare, operator safety and biosecurity;

iii) organise, brief and manage team of people to facilitate humane killing of the relevant animals on the premises in accordance with national regulations and these guidelines;

iv) determine logistics required;

v) monitor operations to ensure animal welfare, operator safety and biosecurity requirements are met;

vi) report upwards on progress and problems;

vii) provide a written report at the conclusion of the killing, describing the practices adopted and their effect on animal welfare, operator safety and biosecurity outcomes.

b) Competencies

i) appreciation of normal animal husbandry practices;

ii) appreciation of animal welfare and the underpinning behavioural, anatomical and physiological processes involved in the killing process;

iii) skills to manage all activities on premises and deliver outcomes on time;

iv) awareness of psychological effects on farmer, team members and general public;

v) effective communication skills.

2. Veterinarian

a) Responsibilities

i) determine and implement the most appropriate killing method to ensure that animals are killed without avoidable pain and distress;

ii) determine and implement the additional requirements for animal welfare, including the order of killing;

iii) ensure that confirmation of animals deaths is carried out by competent persons at appropriate times after the killing procedure;

iv) minimise the risk of disease spread within and from the premises through the supervision of biosecurity procedures;

v) continuously monitor animal welfare and biosecurity procedures;

vi) in cooperation with the leader, prepare a written report at the conclusion of the killing, describing the practices adopted and their effect on animal welfare.
Appendix XXIII (contd)

b) Competencies
   i) ability to assess animal welfare, especially the effectiveness of stunning and killing, and to correct any deficiencies;
   ii) ability to assess biosecurity risks.

3. Animal handlers
   a) Responsibilities
      i) review on-site facilities in terms of their appropriateness;
      ii) design and construct temporary animal handling facilities, when required;
      iii) move and restrain animals.
   b) Competencies
      i) An experience of Animal handling in emergency situations and in close confinement is required;
      ii) an appreciation of biosecurity and containment principles.

4. Slaughterers, Animal killing personnel
   a) Responsibilities
      Humane killing of the animals through effective stunning and killing should be ensured.
   b) Competencies
      i) when required by regulations, licensed to use necessary equipment or licensed to be slaughterers;
      ii) competent to use and maintain relevant equipment;
      iii) competent to use techniques for the species involved;
      iv) competent to assess effective stunning and killing.

5. Carcass disposal personnel
   a) Responsibilities
      An efficient carcass disposal (to ensure killing operations are not hindered) should be ensured.
   b) Competencies
      The personnel should be competent to use and maintain available equipment and apply techniques for the species involved.
6. **Farmer/owner/manager**
   
a) **Responsibilities**
   
i) assist when requested.

b) **Competencies**
   
i) specific knowledge of his/her animals and their environment.

**Article 3.7.6.4.**

### Considerations in planning the humane killing of animals

Many activities will need to be conducted on affected premises, including the humane killing of animals. The team leader should develop a plan for humanely killing animals on the premises which should include consideration of:

1. minimising handling and movement of animals;

2. killing the animals on the affected premises; however, there may be circumstances where the animals may need to be moved to another location for killing; when the killing is conducted at an abattoir, the guidelines in the Chapter on slaughter of animal for human consumption should be followed;

3. the species, number, age and size of animals to be killed, and the order of killing them;

4. methods of killing the animals, and their cost;

5. housing and location of the animals;

6. the availability and effectiveness of equipment needed for killing of the animals;

7. the facilities available on the premises that will assist with the killing;

8. biosecurity and environmental issues;

9. the health and safety of personnel conducting the killing;

10. any legal issues that may be involved, for example where restricted veterinary drugs or poisons may be used, or where the process may impact on the environment; and

11. the presence of other nearby premises holding animals.

In designing a killing plan, it is essential that the method chosen be consistently reliable to ensure that all animals are humanely and quickly killed.
Appendix XXIII (contd)

**Article 3.7.6.5.**

Table summarising killing methods described in Articles 3.7.6.6.-3.7.6.17.

<table>
<thead>
<tr>
<th>Species</th>
<th>Age range</th>
<th>Procedure</th>
<th>Restraint necessary</th>
<th>Animal welfare concerns with inappropriate application</th>
<th>Article reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cattle</td>
<td>all</td>
<td>free bullet</td>
<td>no</td>
<td>non-lethal wounding</td>
<td>3.7.6.6.</td>
</tr>
<tr>
<td></td>
<td>all except neonates</td>
<td>captive bolt - penetrating, followed by pithing or bleeding</td>
<td>yes</td>
<td>ineffective stunning</td>
<td>3.7.6.7.</td>
</tr>
<tr>
<td></td>
<td>adults only</td>
<td>captive bolt - non-penetrating, followed by bleeding</td>
<td>yes</td>
<td>ineffective stunning, regaining of consciousness before killing</td>
<td>3.7.6.8.</td>
</tr>
<tr>
<td></td>
<td>calves only</td>
<td>electrical, two stage application</td>
<td>yes</td>
<td>pain associated with cardiac arrest after ineffective stunning</td>
<td>3.7.6.10.</td>
</tr>
<tr>
<td></td>
<td>calves only</td>
<td>electrical, single application (method 1)</td>
<td>yes</td>
<td>ineffective stunning</td>
<td>3.7.6.11.</td>
</tr>
<tr>
<td></td>
<td>all</td>
<td>injection with barbiturates and other drugs</td>
<td>yes</td>
<td>non-lethal dose, pain associated with injection site</td>
<td>3.7.6.15.</td>
</tr>
<tr>
<td>Sheep and goats</td>
<td>all</td>
<td>free bullet</td>
<td>no</td>
<td>non-lethal wounding</td>
<td>3.7.6.6.</td>
</tr>
<tr>
<td></td>
<td>all except neonates</td>
<td>captive bolt - penetrating, followed by pithing or bleeding</td>
<td>yes</td>
<td>ineffective stunning, regaining of consciousness before killing</td>
<td>3.7.6.7.</td>
</tr>
<tr>
<td></td>
<td>all except neonates</td>
<td>captive bolt - non-penetrating, followed by bleeding</td>
<td>yes</td>
<td>ineffective stunning, regaining of consciousness before killing</td>
<td>3.7.6.8.</td>
</tr>
<tr>
<td></td>
<td>neonates</td>
<td>captive bolt - non-penetrating</td>
<td>yes</td>
<td>non-lethal wounding</td>
<td>3.7.6.8.</td>
</tr>
<tr>
<td></td>
<td>all</td>
<td>electrical, two stage application</td>
<td>yes</td>
<td>pain associated with cardiac arrest after ineffective stunning</td>
<td>3.7.6.10.</td>
</tr>
<tr>
<td></td>
<td>all</td>
<td>electrical, single application (Method 1)</td>
<td>yes</td>
<td>ineffective stunning</td>
<td>3.7.6.11.</td>
</tr>
<tr>
<td></td>
<td>neonates only</td>
<td>CO₂ / air mixture</td>
<td>yes</td>
<td>slow induction of unconsciousness, aversiveness of induction</td>
<td>3.7.6.12.</td>
</tr>
<tr>
<td></td>
<td>neonates only</td>
<td>nitrogen and/or inert gas mixed with CO₂</td>
<td>yes</td>
<td>slow induction of unconsciousness, aversiveness of induction</td>
<td>3.7.6.13.</td>
</tr>
<tr>
<td>Species</td>
<td>Age range</td>
<td>Procedure</td>
<td>Restraint necessary</td>
<td>Animal welfare concerns with inappropriate application</td>
<td>Article reference</td>
</tr>
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<td>----------------------</td>
<td>----------------------------------------------------------------------------</td>
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<td>--------------------------------------------------------------------------------------------------------------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>Sheep and goats</td>
<td>neonates only</td>
<td>nitrogen and/or inert gases</td>
<td>yes</td>
<td>slow induction of unconsciousness, aversiveness of induction</td>
<td>3.7.6.12.</td>
</tr>
<tr>
<td>(contd)</td>
<td>all</td>
<td>injection of barbiturates and other drugs</td>
<td>yes</td>
<td>non-lethal dose, pain associated with injection site</td>
<td>3.7.6.15.</td>
</tr>
<tr>
<td>Pigs</td>
<td>all</td>
<td>free bullet</td>
<td>no</td>
<td>non-lethal wounding</td>
<td>3.7.6.6.</td>
</tr>
<tr>
<td></td>
<td>all except neonates</td>
<td>captive bolt - penetrating, followed by pithing or bleeding</td>
<td>yes</td>
<td>ineffective stunning, regaining of consciousness before death</td>
<td>3.7.6.7.</td>
</tr>
<tr>
<td></td>
<td>neonates only</td>
<td>captive bolt - non-penetrating</td>
<td>yes</td>
<td>non-lethal wounding</td>
<td>3.7.6.8.</td>
</tr>
<tr>
<td></td>
<td>all §</td>
<td>electrical, two stage application</td>
<td>yes</td>
<td>pain associated with cardiac arrest after ineffective stunning</td>
<td>3.7.6.10.</td>
</tr>
<tr>
<td></td>
<td>all</td>
<td>electrical, single application (Method 1)</td>
<td>yes</td>
<td>ineffective stunning</td>
<td>3.7.6.11.</td>
</tr>
<tr>
<td>Poultry</td>
<td>neonates only</td>
<td>CO₂/ air mixture</td>
<td>yes</td>
<td>slow induction of unconsciousness, aversiveness of induction</td>
<td>3.7.6.12.</td>
</tr>
<tr>
<td></td>
<td>neonates only</td>
<td>nitrogen and/or inert gas mixed with CO₂</td>
<td>yes</td>
<td>slow induction of unconsciousness, aversiveness of induction</td>
<td>3.7.6.13.</td>
</tr>
<tr>
<td></td>
<td>neonates only</td>
<td>nitrogen and/or inert gases</td>
<td>yes</td>
<td>slow induction of unconsciousness</td>
<td>3.7.6.14.</td>
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<tr>
<td></td>
<td>all</td>
<td>injection with barbiturates and other drugs</td>
<td>yes</td>
<td>non-lethal dose, pain associated with injection site</td>
<td>3.7.6.15.</td>
</tr>
<tr>
<td>Adults only</td>
<td>captive bolt - non-penetrating</td>
<td>yes</td>
<td>ineffective stunning</td>
<td>3.7.6.8.</td>
<td></td>
</tr>
<tr>
<td>Adults only</td>
<td>maceration</td>
<td></td>
<td>no</td>
<td>non-lethal wounding, non-immediacy;</td>
<td>3.7.6.9.</td>
</tr>
<tr>
<td>Adults only</td>
<td>electrical single application (Method 2)</td>
<td>yes</td>
<td>ineffective stunning</td>
<td>3.7.6.11.</td>
<td></td>
</tr>
<tr>
<td>Adults only</td>
<td>electrical single application, followed by killing (Method 3)</td>
<td>yes</td>
<td>ineffective stunning, regaining of consciousness before killing</td>
<td>3.7.6.11.</td>
<td></td>
</tr>
</tbody>
</table>
### Appendix XXIII (contd)

<table>
<thead>
<tr>
<th>Species</th>
<th>Age range</th>
<th>Procedure</th>
<th>Restraint necessary</th>
<th>Animal welfare concerns with inappropriate application</th>
<th>Article reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poultry (contd)</td>
<td>all</td>
<td>CO₂ / air mixture Method 1 Method 2</td>
<td>yes no</td>
<td>slow induction of unconsciousness, aversiveness of induction</td>
<td>3.7.6.12.</td>
</tr>
<tr>
<td>all</td>
<td>nitrogen and/or inert gas mixed with CO₂</td>
<td>yes</td>
<td>slow induction of unconsciousness, aversiveness of induction</td>
<td>3.7.6.13.</td>
<td></td>
</tr>
<tr>
<td>all</td>
<td>nitrogen and/or inert gases</td>
<td>yes</td>
<td>slow induction of unconsciousness</td>
<td>3.7.6.14.</td>
<td></td>
</tr>
<tr>
<td>all</td>
<td>injection of barbiturates and other drugs</td>
<td>yes</td>
<td>non-lethal dose, pain associated with injection site</td>
<td>3.7.6.15.</td>
<td></td>
</tr>
<tr>
<td>adults only</td>
<td>addition of anaesthetics to feed or water, followed by an appropriate killing method</td>
<td>no</td>
<td>ineffective or slow induction of unconsciousness</td>
<td>3.7.6.16.</td>
<td></td>
</tr>
</tbody>
</table>

* The methods are described in the order of mechanical, electrical and gaseous, not in an order of desirability from an animal welfare viewpoint.

§ The only preclusion against the use of this method for neonates is the design of the stunning tongs that may not facilitate their application across such a small-sized head/body.

#### Free bullet

1. **Introduction**
   
a) A free bullet is a projectile fired from a shotgun, rifle, handgun or purpose-made humane killer.

   b) The most commonly used firearms for close range use are:
      
      i) humane killers (specially manufactured/adapted single-shot weapons);
      
      ii) shotguns (12, 16, 20, 28 bore and .410);
      
      iii) rifles (.22 rimfire);
      
      iv) handguns (various calibres from .32 to .45).

   c) The most commonly used firearms for long range use are rifles (.22, .243, .270 and .308).
d) A free bullet used from long range should be aimed to penetrate the skull or soft tissue at the top of the neck of the animal, to cause irreversible concussion and death and should only be used by properly trained and competent marksmen.

2. Requirements for effective use

a) The marksman should take account of human safety in the area in which he/she is operating. Appropriate vision and hearing protective devices should be worn by all personnel involved.

b) The marksman should ensure that the animal is not moving and in the correct position to enable accurate targeting and the range should be as short as possible (5 – 50 cm for a shotgun) but the barrel should not be in contact with the animal’s head.

c) The correct cartridge, calibre and type of bullet for the different species age and size should be used. Ideally the ammunition should expand upon impact and dissipate its energy within the cranium.

d) Shot animals should be checked to ensure the absence of brain stem reflexes.

**Figure 1.** The optimum shooting position for cattle is at the intersection of two imaginary lines drawn from the rear of the eyes to the opposite horn buds.

![Figure 1](image1)

**Figure 2.** The optimum position for hornless sheep and goats is on the midline just above the eye level, and directing the shot down the line of the spinal cord.

![Figure 2](image2)
Appendix XXIII (contd)

**Figure 3.** The optimum shooting position for heavily horned sheep and horned goats is behind the poll aiming towards the angle of the jaw.

![Diagram of a sheep showing the optimum shooting position.]

**Figure 4.** The optimum shooting position for pigs is just above the eyes level, with the shot directed down the line of the spinal cord.

![Diagram of a pig showing the optimum shooting position.]

3. **Advantages**

   a) Used properly, a free bullet provides a quick and effective method for killing.

   b) It requires minimal or no restraint and can be use to kill from a distance.

   c) It is suitable for killing agitated animals in open spaces.
4. **Disadvantages**
   
a) The method is potentially dangerous to humans and other animals in the area.

b) It has the potential for non-lethal wounding.

c) Destruction of brain tissue may preclude diagnosis of some diseases.

d) Leakage of bodily fluids may present a biosecurity risk.

e) Legal requirements may preclude or restrict use.

f) There is a limited availability of competent personnel.

4. **Conclusions**

   The method is suitable for cattle, sheep, goats and pigs, including large animals in open spaces.

   **Article 3.7.6.7.**

---

**Penetrating captive bolt**

1. **Introduction**

   A penetrating captive bolt is fired from a gun powered by either compressed air or a blank cartridge. There is no free projectile.

   The captive bolt should be aimed on the skull in a position to penetrate the cortex and mid-brain of the animal. The impact of the bolt on the skull produces unconsciousness. Physical damage to the brain caused by penetration of the bolt may result in death, however pithing or bleeding should be performed as soon as possible after the shot to ensure the death of the animal.

2. **Requirements for effective use**

   a) For cartridge powered and compressed air guns, the bolt velocity and the length of the bolt should be appropriate to the species and type of animal, in accordance with the manufacturer’s recommendations.

   b) Captive bolt guns should be frequently cleaned and maintained in good working condition.

   c) More than one gun may be necessary to avoid overheating and a back-up gun should be available in the event of an ineffective shot.

   d) Animals should be restrained; at a minimum they should be penned for cartridge powered guns and in a race for compressed air guns.

   e) The operator should ensure that the animal's head is accessible.

   f) The operator should fire the captive bolt at right angles to the skull in the optimal position (see figures 1, 3 & 4. The optimum shooting position for hornless sheep is on the highest point of the head, on the midline and aim towards the angle of the jaw).
Appendix XXIII (contd)

g) To ensure the death of the animal, pithing or bleeding should be performed as soon as possible after stunning.

h) Animals should be monitored continuously after stunning until death to ensure the absence of brain stem reflexes.

3. Advantages

a) Mobility of cartridge powered equipment reduces the need to move animals.

b) The method induces an immediate onset of a sustained period of unconsciousness.

4. Disadvantages

a) Poor gun maintenance and misfiring, and inaccurate gun positioning and orientation may result in poor animal welfare.

b) Post stun convulsions may make pithing difficult and hazardous.

c) The method is difficult to apply in agitated animals.

d) Repeated use of a cartridge powered gun may result in over-heating.

e) Leakage of bodily fluids may present a biosecurity risk.

f) Destruction of brain tissue may preclude diagnosis of some diseases.

5. Conclusions

The method is suitable for cattle, sheep, goats and pigs (except neonates), when followed by pithing or bleeding.

Article 3.7.6.8.

Captive bolt - non-penetrating

1. Introduction

A non-penetrating captive bolt is fired from a gun powered by either compressed air or a blank cartridge. There is no free projectile.

The gun should be placed on the front of the skull to deliver a percussive blow which produces unconsciousness in cattle (adults only), sheep, goats and pigs, and death in poultry and neonate sheep, goats and pigs. In mammals, bleeding should be performed as soon as possible after the blow to ensure the death of the animal.

2. Requirements for effective use

a) For cartridge powered and compressed air guns, the bolt velocity should be appropriate to the species and type of animal, in accordance with the manufacturer’s recommendations.
b) Captive bolt guns should be frequently cleaned and maintained in good working condition.

c) More than one gun may be necessary to avoid overheating and a back-up gun should be available in the event of an ineffective shot.

d) Animals should be restrained; at a minimum mammals should be penned for cartridge powered guns and in a race for compressed air guns; birds should be restrained in cones, shackles, crushes or by hand.

e) The operator should ensure that the animal’s head is accessible.

f) The operator should fire the captive bolt at right angles to the skull in the optimal position (figures 1-4).

g) To ensure death in non-neonate mammals, bleeding should be performed as soon as possible after stunning.

h) Animals should be monitored continuously after stunning until death to ensure the absence of brain stem reflexes.

3. Advantages

a) The method induces an immediate onset of unconsciousness, and death in birds and neonates.

b) Mobility of equipment reduces the need to move animals

4. Disadvantages

a) As consciousness can be regained quickly in non-neonate mammals, they should be bled as soon as possible after stunning.

b) Laying hens in cages have to be removed from their cages and most birds have to be restrained.

c) Poor gun maintenance and misfiring, and inaccurate gun positioning and orientation may result in poor animal welfare.

d) Post stun convulsions may make bleeding difficult and hazardous.

e) Difficult to apply in agitated animals; such animals may be sedated in advance of the killing procedure.

f) Repeated use of a cartridge powered gun may result in over-heating.

g) Bleeding may present a biosecurity risk.

5. Conclusions

a) The method is suitable for poultry, and neonate sheep, goats and pigs.

b) If bleeding does not present a biosecurity issue, this is a suitable method for cattle (adults only), and non-neonate sheep, goats and pigs when followed by bleeding.
Appendix XXIII (contd)

Article 3.7.6.9.

Maceration

1. **Introduction**

Maceration, utilising a mechanical apparatus with rotating blades or projections, causes immediate fragmentation and death in day-old poultry and embryonated eggs.

2. **Requirements**

   a) Maceration requires specialised equipment which should be kept in excellent working order.

   b) The rate of introducing the birds should not allow the equipment to jam, birds to rebound from the blades or the birds to suffocate before they are macerated.

3. **Advantages**

   a) Procedure results in immediate death.

   b) Large numbers can be killed quickly.

4. **Disadvantages**

   a) Specialised equipment is required.

   b) Macerated tissues may present a biosecurity issue.

5. **Conclusion**

   The method is suitable for killing day-old poultry and embryonated eggs.

Article 3.7.6.10.

Electrical – two stage application

1. **Introduction**

A two stage application of electric current comprises firstly an application of current to the head by scissor-type tongs, immediately followed by an application of the tongs across the chest in a position that spans the heart.

The application of sufficient electric current to the head will induce ‘tonic/ clonic’ epilepsy and unconsciousness. Once the animal is unconscious, the second stage will induce ventricular fibrillation (cardiac arrest) resulting in death. The second stage (the application of low frequency current across the chest) should only be applied to unconscious animals to prevent unacceptable levels of pain.

Figure 6. Scissor-type stunning tongs.
2. Requirements for effective use

a) The stunner control device should generate a low frequency (30 – 60 Hz) current with a minimum voltage of 250 volts true RMS under load.

b) Appropriate protective clothing (including rubber gloves and boots) should be worn.

c) Animals should be restrained, at a minimum free-standing in a pen, close to an electrical supply.

d) Two team members are required, the first to apply the electrodes and the second to manipulate the position of the animal to allow the second application to be made.

e) A stunning current should be applied via scissor-type stunning tongs in a position that spans the brain for a minimum of 3 seconds; immediately following the application to the head, the electrodes should be transferred to a position that spans the heart and the electrodes applied for a minimum of 3 seconds.

f) Electrodes should be cleaned regularly and after use, to enable optimum electrical contact to be maintained.

g) Animals should be monitored continuously after stunning until death to ensure the absence of brain stem reflexes.

3. Advantages

a) The application of the second stage minimises post-stun convulsions and therefore the method is particularly effective with pigs.

b) Non-invasive technique minimises biosecurity risk.

4. Disadvantages

a) The method requires a reliable supply of electricity.

b) The electrodes must be applied and maintained in the correct positions to produce an effective stun and kill.

c) Most stunner control devices utilise low voltage impedance sensing as an electronic switch prior to the application of high voltages; in unshorn sheep, contact impedance may be too high to switch on the required high voltage (especially during stage two).

d) The procedure may be physically demanding, leading to operator fatigue and poor electrode placement.

5. Conclusion

The method is suitable for calves, sheep and goats, and especially for pigs (over one week of age).

Article 3.7.6.11.

Electrical – single application

1. Method 1

Method 1 comprises the single application of sufficient electrical current to the head and back, to simultaneously stun the animal and fibrillate the heart. Provided sufficient current is applied in a position that spans both the brain and heart, the animal will not recover consciousness.
Appendix XXIII (contd)

a) Requirements for effective use
   
i) The stunner control device should generate a low frequency (30 – 60 Hz) current with a minimum voltage of 250 volts true RMS under load.
   
ii) Appropriate protective clothing (including rubber gloves and boots) should be worn.
   
iii) Animals should be individually and mechanically restrained close to an electrical supply as the maintenance of physical contact between the stunning electrodes and the animal is necessary for effective use.
   
iv) The rear electrode should be applied to the back, above or behind the heart, and then the front electrode in a position that is forward of the eyes, with current applied for a minimum of 3 seconds.
   
v) Electrodes should be cleaned regularly between animals and after use, to enable optimum electrical contact to be maintained.
   
vi) Water or saline may be necessary to improve electrical contact with sheep.
   
vii) An effective stun and kill should be verified by the absence of brain stem reflexes.

b) Advantages
   
i) Method 1 stuns and kills simultaneously.
   
ii) It minimises post-stun convulsions and therefore is particularly effective with pigs.
   
iii) A single team member only is required for the application.
   
iv) Non-invasive technique minimises biosecurity risk.

c) Disadvantages
   
i) Method 1 requires individual mechanical animal restraint.
   
ii) The electrodes must be applied and maintained in the correct positions to produce an effective stun and kill.
   
iii) Method 1 requires a reliable supply of electricity.

d) Conclusion

Method 1 is suitable for calves, sheep, goats, and pigs (over 1 week of age).

2. Method 2

Method 2 stuns and kills by drawing inverted and shackled poultry through an electrified waterbath stunner. Electrical contact is made between the ‘live’ water and earthed shackle and, when sufficient current is applied, poultry will be simultaneously stunned and killed.
a) Requirements for effective use

i) A mobile waterbath stunner and a short loop of processing line are required.

ii) A low frequency (30-60 Hz) current applied for a minimum of 3 seconds is necessary to stun and kill the birds.

iii) Poultry need to be manually removed from their cage, house or yard, inverted and shackled onto a line which conveys them through a waterbath stunner with their heads fully immersed.

iv) The required minimum currents to stun and kill dry birds are:
   - Quail - 100 mA/bird
   - Chickens – 160 mA/bird
   - Ducks & Geese – 200 mA/bird
   - Turkeys – 250 mA/bird.
   A higher current is required for wet birds.

v) An effective stun and kill should be verified by the absence of brain stem reflexes.

b) Advantages

i) Method 2 stuns and kills simultaneously.

ii) It is capable of processing large numbers of birds reliably and effectively.

iii) This non-invasive technique minimises biosecurity risk.

c) Disadvantages

i) Method 2 requires a reliable supply of electricity.

ii) Handling, inversion and shackling of birds are required.

d) Conclusion

Method 2 is suitable for large numbers of poultry.

3. Method 3

Method 3 comprises the single application of sufficient electrical current to the head of poultry in a position that spans the brain, causing unconsciousness; this is followed by a killing method (Article 17).

a) Requirements for effective use

i) The stunner control device should generate sufficient current (more than 300 mA/bird) to stun.

ii) Appropriate protective clothing (including rubber gloves and boots) should be worn.
Appendix XXIII (contd)

iii) Birds should be restrained, at a minimum manually, close to an electrical supply.

iv) A stunning current should be applied in a position that spans the brain for a minimum of 3 seconds; immediately following this application, the birds should be killed (Article 17).

v) Electrodes should be cleaned regularly and after use, to enable optimum electrical contact to be maintained.

vi) Birds should be monitored continuously after stunning until death to ensure the absence of brain stem reflexes.

b) Advantages

Non-invasive technique (when combined with neck cervical dislocation) minimises biosecurity risk.

c) Disadvantages

i) Method 3 requires a reliable supply of electricity.

ii) The electrodes must be applied and maintained in the correct position to produce an effective stun.

iii) Birds must be individually restrained.

iv) It must be followed by a killing method.

d) Conclusion

Method 3 is suitable for small numbers of poultry.

Article 3.7.6.12.

CO₂ / air mixture

1. Introduction

Controlled atmosphere killing is performed by exposing animals to a predetermined gas mixture, either by placing them in a gas-filled container or apparatus (Method 1) or by the gas being introduced into a poultry house (Method 2).

Inhalation of carbon dioxide (CO₂) induces respiratory and metabolic acidosis and hence reduces the pH of cerebrospinal fluid (CSF) and neurones thereby causing unconsciousness and, after prolonged exposure, death.
2. **Method 1**

The animals are placed in a gas-filled container or apparatus.

a) Requirements for effective use in a container or apparatus
   
i) Containers or apparatus should allow the required gas concentration to be maintained and accurately measured.
   
ii) When animals are exposed to the gas individually or in small groups in a container or apparatus, the equipment used should be designed, constructed, and maintained in such a way as to avoid injury to the animals and allow them to be observed.
   
iii) Animals should be introduced into the container or apparatus after it has been filled with the required CO₂ concentration, and held in this atmosphere until death is confirmed.
   
iv) Team members should ensure that there is sufficient time allowed for each batch of animals to die before subsequent ones are introduced into the container or apparatus.
   
v) Containers or apparatus should not be overcrowded and measures are needed to avoid animals suffocating by climbing on top of each other.

b) Advantages
   
i) CO₂ is readily available.
   
ii) Application methods are simple.

c) Disadvantages
   
i) The need for properly designed container or apparatus special equipment
   
ii) The aversive nature of high CO₂ concentrations
   
iii) No immediate loss of consciousness
   
iv) The risk of suffocation due to overcrowding
   
v) Difficulty in verifying death while the animals are in the container or apparatus.

d) Conclusion
   
Method 1 is suitable for use in poultry and neonatal sheep, goats and pigs.

3. **Method 2**

The gas is introduced into a poultry house.

a) Requirements for effective use in a poultry house
   
i) Prior to introduction of the CO₂, the poultry house should be appropriately sealed to allow control over the gas concentration.
Appendix XXIII (contd)

ii) The house should be gradually filled with CO$_2$ so that all birds are exposed to a concentration of >40% until they are dead; a vaporiser may be required to prevent freezing.

iii) Devices should be used to accurately measure the gas concentration at the highest level of birds.

b) Advantages

i) Applying gas to birds in situ eliminates the need to manually remove live birds.

ii) CO$_2$ is readily available.

iii) Gradual raising of CO$_2$ concentration minimises the aversiveness of the induction of unconsciousness.

c) Disadvantages

i) It is difficult to determine volume of gas required to achieve adequate concentrations of CO$_2$ in some poultry houses.

ii) It is difficult to verify death while the birds are in the poultry house.

d) Conclusion

Method 2 is suitable for use in poultry in closed-environment sheds

Article 3.7.6.13.

Nitrogen and/or inert gas mixed with CO$_2$

1. Introduction

CO$_2$ may be mixed in various proportions with nitrogen or an inert gas eg argon, and the inhalation of such mixtures leads to hypercapnic-hypoxia and death when the oxygen concentration by volume is $\leq$2%. This method involves the introduction of animals into a container or apparatus containing the gases. Such mixtures do not induce immediate loss of consciousness, therefore the aversiveness of various gas mixtures containing high concentrations of CO$_2$ and the respiratory distress occurring during the induction phase, are important animal welfare considerations.

Pigs and poultry appear not to find low concentrations of CO$_2$ strongly aversive, and a mixture of nitrogen or argon with $\leq$30% CO$_2$ by volume and $\leq$2% O$_2$ by volume can be used for killing poultry and neonatal sheep, goats and pigs.

2. Requirements for effective use

a) Containers or apparatus should allow the required gas concentrations to be maintained, and the O$_2$ and CO$_2$ concentrations accurately measured during the killing procedure.
b) When animals are exposed to the gases individually or in small groups in a container or apparatus, the equipment used should be designed, constructed, and maintained in such a way as to avoid injury to the animals and allow them to be observed.

c) Animals should be introduced into the container or apparatus after it has been filled with the required gas concentrations (with $\leq 2\%$ $O_2$), and held in this atmosphere until death is confirmed.

d) Team members should ensure that there is sufficient time allowed for each batch of animals to die before subsequent ones are introduced into the container or apparatus.

e) Containers or apparatus should not be overcrowded and measures are needed to avoid animals suffocating by climbing on top of each other.

3. Advantages

Low concentrations of CO$_2$ cause little aversiveness and, in combination with nitrogen or an inert gas, produces a fast induction of unconsciousness.

4. Disadvantages

a) A properly designed container or apparatus is needed.

b) It is difficult to verify death while the animals are in the container or apparatus.

c) There is no immediate loss of consciousness.

d) Exposure times required to kill are considerable.

5. Conclusion

The method is suitable for poultry and neonatal sheep, goats and pigs.

Article 3.7.6.14.

Nitrogen and/or inert gasses

1. Introduction

This method involves the introduction of animals into a container or apparatus containing nitrogen or an inert gas such as argon. The controlled atmosphere produces leads to unconsciousness and death from hypoxia.

Research has shown that hypoxia is not aversive to pigs and poultry, and it doesn't induce any signs of respiratory distress prior to loss of consciousness.

2. Requirements for effective use

a) Containers or apparatus should allow the required gas concentrations to be maintained, and the $O_2$ concentration accurately measured.
Appendix XXIII (contd)

b) When animals are exposed to the gases individually or in small groups in a container or apparatus, the equipment used should be designed, constructed, and maintained in such a way as to avoid injury to the animals and allow them to be observed.

c) Animals should be introduced into the container or apparatus after it has been filled with the required gas concentrations (with $\leq 2\%$ O$_2$), and held in this atmosphere until death is confirmed.

d) Team members should ensure that there is sufficient time allowed for each batch of animals to die before subsequent ones are introduced into the container or apparatus.

e) Containers or apparatus should not be overcrowded and measures are needed to avoid animals suffocating by climbing on top of each other.

3. Advantages

Animals are unable to detect nitrogen or inert gases, and the induction of hypoxia by this method is not aversive to animals.

4. Disadvantages

a) A properly designed container or apparatus is needed.

b) It is difficult to verify death while the animals are in the container or apparatus.

c) There is no immediate loss of consciousness.

d) Exposure times required to kill are considerable.

5. Conclusion

The method is suitable for poultry and neonatal sheep, goats and pigs.

Article 3.7.6.15.

Lethal injection

1. Introduction

A lethal injection using high doses of anaesthetic and sedative drugs causes CNS depression, unconsciousness and death. In practice, barbiturates in combination with other drugs are commonly used.

2. Requirements for effective use

a) Doses and routes of administration that cause rapid loss of consciousness followed by death should be used.

b) Prior sedation may be necessary for some animals.

c) Intravenous administration is preferred, but intraperitoneal or intramuscular administration may be appropriate, especially if the agent is non-irritating.
Appendix XXIII (contd)

d) Animals should be restrained to allow effective administration.

e) Animals should be monitored to ensure the absence of brain stem reflexes.

3. Advantages

a) The method can be used in all species.

b) Death can be induced smoothly.

4. Disadvantages

a) Restraint and/or sedation may be necessary prior to injection.

b) Some combinations of drug type and route of administration may be painful, and should only be used in unconscious animals.

c) Legal requirements may restrict use to veterinarians.

d) Contaminated carcasses may present a risk to other wild or domestic animals.

5. Conclusion

The method is suitable for killing small numbers of cattle, sheep, goats, pigs and poultry.

Article 3.7.6.16.

Addition of anaesthetics to feed or water

1. Introduction

An anaesthetic agent which can be mixed with poultry feed or water may be used to kill poultry in houses. Poultry which are only anaesthetised need to be killed by another method such as cervical dislocation.

2. Requirements for effective use

a) Sufficient quantities of anaesthetic need to be ingested rapidly for effective response.

b) Intake of sufficient quantities is facilitated if the birds are fasted or water is withheld.

c) Must be followed by killing (see Article 3.7.6.17) if birds are anaesthetised only.

3. Advantages

a) Handling is not required until birds are anaesthetised.

b) There may be biosecurity advantages in the case of large numbers of diseased birds.

4. Disadvantages

a) Non-target animals may accidentally access the medicated feed or water when provided in an open environment.
Appendix XXIII (contd)

b) Dose taken is unable to be regulated and variable results may be obtained.

c) Animals may reject adulterated feed or water due to illness or adverse flavour.

d) The method may need to be followed by killing.

e) Care is essential in the preparation and provision of treated feed or water, and in the disposal of uneaten treated feed/water and contaminated carcasses.

5. Conclusion

The method is suitable for killing large numbers of poultry in houses.

Article 3.7.6.17.

Killing methods in unconscious animals

1. Method 1: Cervical dislocation (manual and mechanical)

a) Introduction

Poultry may be killed by either manual cervical dislocation (stretching) or mechanical neck crushing with a pair of pliers. Both methods result in death from asphyxiation and/or cerebral anoxia.

b) Requirements for effective use

i) Killing should be performed either by manually or mechanically stretching the neck to sever the spinal cord or by using mechanical pliers to crush the cervical vertebrae with consequent major damage to the spinal cord.

ii) Consistent results require strength and skill so team members should be rested regularly to ensure consistently reliable results.

iii) Birds should be monitored continuously until death to ensure the absence of brain stem reflexes.

c) Advantages

i) It is a non-invasive killing method.

ii) It can be performed manually on small birds.

d) Disadvantages

i) Operator fatigue.

ii) The method is more difficult in larger birds.

e) Conclusion

This method is suitable for killing unconscious poultry.
2. **Method 2: Decapitation**
   a) **Introduction**
   Decapitation results in death by cerebral ischaemia using a guillotine or knife.
   b) **Requirements for effective use**
   The required equipment should be kept in good working order.
   c) **Advantages**
   The technique is effective and does not require monitoring.
   d) **Disadvantages**
   The working area is contaminated with body fluids.
   e) **Conclusion**
   This method is suitable for killing unconscious poultry.

3. **Method 3: Pithing**
   a) **Introduction**
   Pithing is a method of killing animals which have been stunned by a penetrating captive bolt, without immediate death. Pithing results in the physical destruction of the brain and upper regions of the spinal cord, through the insertion of a rod or cane through the bolt hole.
   b) **Requirements for effective use**
   i) Pithing cane or rod is required.
   ii) An access to the head of the animal and to the brain through the skull is required.
   iii) Animals should be monitored continuously until death to ensure the absence of brain stem reflexes.
   c) **Advantages**
   The technique is effective in producing immediate death.
   d) **Disadvantages**
   i) A delayed and/or ineffective pithing due to convulsions may occur.
   ii) The working area is contaminated with body fluids.
Appendix XXIII (contd)

e) Conclusion

This method is suitable for killing unconscious animals which have been stunned by a
penetrating captive bolt.

4. Method 4: Bleeding

a) Introduction

Bleeding is a method of killing animals through the severance of the major blood vessels in the
neck or chest that results in a rapid fall in blood pressure, leading to cerebral ischaemia and
death.

b) Requirements for effective use

i) A sharp knife is required.

ii) An access to the neck or chest of the animal is required.

iii) Animals should be monitored continuously until death to ensure the absence of brain stem
reflexes.

c) Advantages

The technique is effective in producing death after an effective stunning method which does not
permit pithing.

d) Disadvantages

a) A delayed and/or ineffective bleeding due to convulsions may occur.

b) The working area is contaminated with body fluids.

e) Conclusion

This method is suitable for killing unconscious animals.
Appendix XXIV

APPENDIX X.X.X.

GUIDELINES FOR THE CONTROL OF HAZARDS OF ANIMAL HEALTH AND PUBLIC HEALTH IMPORTANCE THROUGH ANTE- AND POST-MORTEM MEAT INSPECTION

Introduction

Foodborne disease and zoonoses are important public health problems and important causes of decreased economic productivity in developed and developing countries. Similarly, transmission of hazards of animal health importance via the food chain and associated by-products can result in significant economic loss in livestock. Inspection of animals at slaughter can provide a valuable contribution to surveillance for certain diseases of animal and public health importance. Control and/or reduction of hazards of animal and public health importance by ante- and post-mortem meat inspection are a core responsibility of Veterinary Services.

Purpose

These guidelines provide a basis for future development of OIE standards for animal production food safety.

Hygienic practice throughout the food chain

The Codex Alimentarius Code of Hygienic Practice for Meat 50 (CHPM) constitutes the primary international standard for meat hygiene and incorporates a risk-based approach to application of sanitary measures throughout the food chain. Ante-mortem inspection is described as a primary component of meat hygiene pre-slaughter, and post-mortem inspection is described as a primary component of process control in post-slaughter meat hygiene. The CHPM specifically recognises the dual objectives that slaughterhouse inspection activities deliver in terms of animal and public health.

The CHPM does not provide inspection measures for specific hazards or organoleptically detected abnormalities, which remain the responsibility of national competent authorities. The animal and public health risks associated with livestock populations vary across regions and animal husbandry systems, and ante- and post-mortem inspection needs to be tailored to the individual country situation and its animal and public health objectives.

The CHPM provides a platform for development of meat hygiene systems that are based on risk assessment. There are few risk assessment models or relevant scientific information available on public health hazards, making difficult the development of risk-based standards for food-borne zoonoses. While this scientific information is being accumulated, ante- and post-mortem inspection systems will remain dependent on traditional approaches.

50 Code of Hygienic Practice for Meat, CAC/RCP 58-2005
Veterinary Services and meat inspection programmes

Veterinary Services are primarily responsible for the development of ante- and post-mortem meat inspection programmes. Wherever possible, inspection procedures should be risk-based and management systems should reflect international norms. In respect of ante- and post-mortem inspection as a component of meat hygiene, responsibilities of Veterinary Services include:

- Risk assessment
- Establishment of policies and standards
- Design and management of inspection programmes
- Assurance and certification of appropriate delivery of inspection and compliance activities
- Dissemination of information throughout the food chain

Risk assessment

Veterinary Services should utilise risk assessment to the greatest extent possible in the development of sanitary measures. Veterinary Services should give priority to addressing microbiological contamination, rather than gross abnormalities detected at ante and post-mortem inspection, as this has been found to be the most important source of hazards.

Microbiological, serological or other testing at single-animal and herd level as part of ante- and post-mortem inspection should be used to support surveillance, as well as risk assessment of prioritised foodborne hazards. The information gathered should be linked to human disease data to allow an assessment of the effectiveness of various management options, as well as a general evaluation of food sources of foodborne disease.

Application of a generic framework should provide a systematic and consistent process for managing all biosecurity risks, while accommodating the different risk assessment methodologies used in animal and public health.

Establishment of policies and standards

The national competent authority(s) should provide an appropriate institutional environment to allow Veterinary Services to develop the necessary policies and standards.

As well as meeting public health objectives, policies and standards relating to ante- and post-mortem inspection should aim to detect and remove hazards of animal health significance from the food chain. This may be achieved by the removal of live animals at ante-mortem inspection or by the removal of specific tissues at post-mortem inspection.

Veterinary Services should integrate their activities to the maximum extent possible and practicable so as to increase the efficacy of policies to prevent duplication of effort and unnecessary costs e.g. within the process of international certification.

Design and management of inspection programmes

In meeting animal and public health objectives prescribed in national legislation or required by importing countries, Veterinary Services contribute through the direct performance of some veterinary tasks or through the auditing of animal and public health activities conducted by other agencies or the private sector. To this end, Veterinary Services provide assurances domestically and to trading partners that safety and suitability standards have been met.
Veterinary Services should allow flexibility in meat inspection service delivery through an officially recognised competent body operating under its supervision and control. In recognition of the contribution of industry to food safety, quality assurance systems may be extended in the case of ante- and post-mortem inspection to systems that integrate industry and Veterinary Services activities. Nevertheless, Veterinary Services should take into account the factors identified in Chapter 1.3.3 on the Evaluation of Veterinary Services. For example, 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 Services, the Veterinary Services should specify the competency requirements for all such persons and verify their performance.

Assurance and certification

Assurance and certification of appropriate delivery of inspection and compliance activities is a vital function of Veterinary Services. International health certificates providing official assurances for trading of meat must engender full confidence to the country of importation.

Dissemination of information

Organisation and dissemination of information throughout the food chain involves multidisciplinary inputs. To ensure the effective implementation of ante- and post-mortem inspection procedures, Veterinary Services should have in place systems for the monitoring of these procedures and the exchange of information gained. Animal identification and 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 to processors.
**CHAPTER 1.3.7.**

**ANIMAL IDENTIFICATION AND TRACEABILITY**

**Proposed definitions** (to be located in Chapter 1.1.1)

*Animal identification* means the identification and registration of an animal individually or collectively by its *epidemiological unit* or group. Methods of animal identification include tag, brand, tattoo, transponder (microchip), collar, ring and mark.

*Animal identification system* means the inclusion and linking of components such as *identification of establishments/owners*, the person(s) responsible for the animal(s) and records with *animal identification*.

*Animal traceability* means the ability to follow an animal during specified stage(s) of its life.

*Individual identification* means the identification of each animal using a unique identifier.

*Group identification* means the identification of a group of animals using a unique group identifier.

*Register* means the system by which animal identification and traceability information is securely stored and appropriately accessed by the *Competent Authority*.

**General principles**

1. There is a critical relationship between *animal identification* and the traceability of animals and products of animal origin.

2. *Animal traceability* and traceability of products of animal origin should have the capability to be linked to food product traceability in order to maintain traceability throughout the food chain.

3. *Animal identification* and *animal traceability* are important tools for addressing animal health (including zoonoses) and food safety, and may significantly improve the effectiveness of the management of disease outbreaks and food safety incidents, vaccination programmes, herd/flock husbandry, zoning/compartmentalisation, surveillance, early response and notification systems, animal movement controls and assurances of safety in trade.

4. The objective(s) of *animal identification* and *animal traceability* for a particular country, zone or compartment, and the approach used, should be clearly defined, following an assessment of the risks to be addressed, and a consideration of the factors listed below. They should be defined through consultation between the *Veterinary Administration* and relevant sector(s)/stakeholders prior to implementation, and periodically reviewed.

5. There are various factors which may determine the chosen approach. Factors such as the outcomes of the risk assessment, the animal health situation (including zoonoses), animal population parameters (such as species and breeds, numbers and distribution), types of production, animal movement patterns, available technologies, trade in animals and animal products, cost/benefit analysis and other economic considerations, and cultural aspects, should be taken into account when designing the approach. Whatever approach is used, it should comply with relevant OIE standards to ensure that the defined objectives are able to be achieved.
6. *Animal identification* and *animal traceability* should be under the responsibility of the *Veterinary Administration*.

7. The *Veterinary Administration*, in consultation with relevant governmental agencies and the private sector, should establish a legal framework for the implementation and enforcement of *animal identification* and *animal traceability* in the country. In order to facilitate compatibility and consistency, relevant international standards and obligations should be taken into account. This legal framework should include elements such as the objectives, scope, organisational arrangements including the choice of technologies used for identification and registration, obligation of the parties, confidentiality, accessibility issues and the efficient exchange of information.
CHAPTER 2.5.4.

EQUINE INFECTIOUS ANAEMIA

Article 2.5.4.1.

Standards for diagnostic tests are described in the *Terrestrial Manual*.

Article 2.5.4.2.

*Veterinary Administrations of importing countries* should require:

for equines imported on a permanent basis

the presentation of an *international veterinary certificate* attesting that:

1. the animals showed no clinical sign of equine infectious anaemia (EIA) on the day of shipment and during the 48 hours prior to shipment;

2. for breeding animals only, no *case* of EIA has been associated with any premises where the animals were kept during the 3 months prior to shipment;

3. the animals were subjected to a diagnostic test for EIA with negative results during the 30 days prior to shipment.

Article 2.5.4.3.

*Veterinary Administrations of importing countries* should require:

for equines imported on a temporary basis

the presentation of an *international veterinary certificate* attesting that:

4. the animals showed no clinical sign of EIA on the day of shipment and during the 48 hours prior to shipment;

5. no *case* of EIA has been associated with any premises where the animals were kept during the 3 months prior to shipment;

3. the animals were subjected to a diagnostic test for EIA with negative results during the 30 days prior to shipment (the negative response to the serological test remains valid for 120 days).

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CHAPTER 2.5.6.

EQUINE PIROPLASMOSIS

Article 2.5.6.1.
Standards for diagnostic tests are described in the Terrestrial Manual.

Article 2.5.6.2.

Veterinary Administrations of importing countries should require:

for equines

the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of equine piroplasmosis on the day of shipment;
2. were subjected to diagnostic tests for equine piroplasmosis (\textit{Babesia Theileria equi} and \textit{Babesia caballi}) with negative results during the 30 days prior to shipment;
3. were maintained free from ticks during the 30 days prior to shipment, treated against ticks within the 7 days prior to shipment (the importing country may decide to import only during seasons when ticks are not active on its territory).

Article 2.5.6.3.

Veterinary Administrations of importing countries should consider the possibility of importing competition horses on a temporary basis and which are positive to the testing procedure referred to in point 2) of Article 2.5.6.2. under the following safeguards:

1. the horses are accompanied by a passport in conformity with the model contained in Appendix 4.1.5.;
2. the Veterinary Administrations of importing countries require the presentation of an international veterinary certificate attesting that the animals:
   a) showed no clinical sign of equine piroplasmosis on the day of shipment;
   b) were treated against ticks within the 7 days prior to shipment;
3. the horses are kept in an area where necessary precautions are taken to control ticks and that is under the direct supervision of the Veterinary Authority;
4. the horses are regularly examined for the presence of ticks under the direct supervision of the Veterinary Authority.

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*OIE Terrestrial Animal Health Standards Commission/September 2005*
CHAPTER 2.5.7.

EQUINE RHINOPNEUMONITIS

Article 2.5.7.1.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 2.5.7.2.

Veterinary Administrations of importing countries should require:

for equines

the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of equine rhinopneumonitis on the day of shipment and during the 21 days 3 months prior to shipment;

2. were kept for the 21 days 3 months prior to shipment in an establishment where no case of equine rhinopneumonitis was officially reported during that period.

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CHAPTER 2.5.8.

GLANDERS

Article 2.5.8.1.

For the purposes of this Terrestrial Code, the incubation period for glanders shall be 6 months.

Standards for diagnostic tests are described in the Terrestrial Manual.

Article 2.5.8.2.

Glanders free country

A country may be considered free from glanders when:

1. glanders is notifiable in the country;
2. no case of glanders has been reported during confirmed for at least the past 3 last 2 years.

When importing equines for immediate slaughter from an infected country (see Article 2.5.8.5.), a glanders free country will not be considered as infected if one of the imported equines is found infected.

The conditions for such imports will require direct transport of the animals from the place of disembarkation to a designated abattoir and completion of cleaning and disinfection of the means of transport, the lairages and the abattoir immediately after use. These conditions should be prescribed and enforced by the Veterinary Administration.

Article 2.5.8.3.

When importing from glanders free countries, Veterinary Administrations should require:

for equines

the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical signs evidence of glanders on the day of shipment;
2. were kept since birth, or for the past 6 months prior to shipment, in the exporting country; or
3. were subjected to a test as prescribed in the Terrestrial Manual the mallein test and/or the complement fixation test for glanders with negative results, during the 15 days prior to shipment.

Article 2.5.8.4.

When importing from countries considered infected with glanders, Veterinary Administrations should require:

for equines

the presentation of an international veterinary certificate attesting that the animals:

1. showed no clinical sign of glanders on the day of shipment;
Appendix XXIX (contd)

2. were kept for the 6 months prior to shipment in an establishment where no case of glanders was officially reported during that period;

3. were subjected to a test as prescribed in the Terrestrial Manual, the mallein test and the complement fixation test for glanders with negative results, during the 15 days prior to shipment.

Article 2.5.8.5.

When importing from countries considered infected with glanders, Veterinary Administrations should require:

for equines for immediate slaughter

the presentation of an international veterinary certificate attesting that the animals showed no clinical sign of glanders on the day of shipment. (See also Article 2.5.8.2.)
CHAPTER 2.5.10.
EQUINE VIRAL ARTERITIS

Article 2.5.10.1.

The infective period for equine viral arteritis (EVA) shall be 28 days for mares, and geldings, and sexually immature equines. The health status of seropositive stallions should be checked to ensure that they do not shed equine arteritis virus in their semen.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 2.5.10.2.

Veterinary Administrations of importing countries should require:

for uncastrated male equines imported on a temporary basis for breeding or on a permanent basis

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;

2. were subjected to two tests for EVA as prescribed in the Terrestrial Manual diagnostic on blood samples at least 14 days apart with negative results, during the 28 days prior to shipment; or

3. were subjected between 6 and 12 months of age to a diagnostic test for EVA as prescribed in the Terrestrial Manual on a blood sample with negative results, immediately vaccinated for EVA and regularly revaccinated; or

4. have been subjected to a diagnostic 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 shipment which were subjected to two tests for EVA as prescribed in the Terrestrial Manual diagnostic 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 virus isolation test for EVA as prescribed in the Terrestrial Manual with negative results (under study), carried out on semen collected during the 28 days prior to shipment.

Article 2.5.10.3.

Veterinary Administrations of importing countries should require:

for uncastrated male equines imported on a temporary basis other than for breeding, and for equines other than uncastrated males

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;
2. were subjected, during the 28 days prior to shipment, to two diagnostic tests for EVA as prescribed in the *Terrestrial Manual* on blood samples collected at least 14 days apart, which demonstrated negative results or a stable or declining antibody titre;

3. were subjected, between 6 and 12 months of age, to a diagnostic test for EVA as prescribed in the *Terrestrial Manual* on a blood sample, with negative results, and immediately vaccinated for EVA and regularly revaccinated.

**Article 2.5.10.4.**

_Veterinary Administrations of importing countries_ should require:

_for fresh semen_

the presentation of an *international veterinary certificate* attesting that the donor animals:

1. were kept for the 28 to 30 days prior to semen collection in an *establishment* where no equine has shown any clinical sign of EVA during that period;

2. showed no clinical sign of EVA on the day of semen collection;

3. were subjected between 6 and 12 months of age to a diagnostic test for EVA as prescribed in the *Terrestrial Manual* on a blood sample with negative results, and immediately vaccinated for EVA and regularly revaccinated; or

4. were subjected to a diagnostic 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 not been used for natural breeding from the time of the taking of the blood sample to the time of semen collection; or

5. were subjected to a diagnostic test for EVA as prescribed in the *Terrestrial Manual* on a blood sample with positive results and then: either
   
   a) were test mated, within 12 months one year prior to semen collection, to two mares which showed negative results to two diagnostic tests as prescribed in the *Terrestrial Manual* on blood samples collected at the time of test mating and again 28 days after the test mating, or
   
   b) were subjected to a *virus isolation test* as prescribed in the *Terrestrial Manual* with negative results (under study), carried out on semen collected within one year prior to collection of the semen to be exported.

**Article 2.5.10.5.**

_Veterinary Administrations of importing countries_ should require:

_for frozen semen_

the presentation of an *international veterinary certificate* attesting that the donor animals:

1. showed no clinical sign of EVA on the day of semen collection;

2. were subjected to a diagnostic test for EVA as prescribed in the *Terrestrial Manual* on a blood sample with negative results not less than 14 days after semen collection; or
3. were subjected, between 6 and 12 months of age, to a diagnostic test for EVA as prescribed in the *Terrestrial Manual* on a blood sample with negative results, and immediately vaccinated for EVA and regularly revaccinated; or

4. were subjected to a diagnostic test for EVA as prescribed in the *Terrestrial Manual* on a blood sample with positive results and then: either

a) were test mated, within 12 months one year prior to or as soon as possible after semen collection, to two mares which showed negative results to two diagnostic tests as prescribed in the *Terrestrial Manual* on blood samples collected at the time of test mating and again 28 days after the test mating, or

b) were subjected to a virus isolation test as prescribed in the *Terrestrial Manual* with negative results (under study), carried out on semen collected within one year prior to collection of the semen to be exported.
CHAPTER 2.X.X.
AFRICAN HORSE SICKNESS

Article 2.x.x.1.
For the purposes of this Terrestrial Code, the infective period for African horse sickness (AHS) shall be 40 days for domestic horses.

All countries or zones adjacent to a country or zone not having free status should determine their AHS status from an ongoing surveillance programme (in accordance with Appendix 3.8.X.). 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 AHS.

Standards for diagnostic tests and vaccines are described in the Terrestrial Manual.

Article 2.x.x.2.
AHS free country or zone
1. A country or a zone may be considered free from AHS when the disease is notifiable in the whole country and either:
   a) the country or zone is not adjacent to a country or zone not having a free status; or
   b) historical freedom as described in Appendix 3.8.1. has demonstrated no evidence of AHS in the country or zone, or
   c) a surveillance programme as described in Appendix 3.8.X. has demonstrated no evidence of AHS in the country or zone during the past 2 years, including in wildlife; or
   d) a surveillance programme has demonstrated no evidence of Culicoides likely to be competent AHS vectors in the country or zone.

2. An AHS free country or zone in which surveillance has found no evidence that Culicoides likely to be competent AHS vectors are present will not lose its free status through the importation of vaccinated or seropositive animals, semen or embryos from infected countries or zones.

3. An AHS free country or zone in which surveillance has found evidence that Culicoides likely to be competent AHS vectors are present will not lose its free status through the importation of vaccinated or seropositive domestic horses from infected countries or zones, provided:
   a) the animals have been vaccinated, in accordance with the Terrestrial Manual, at least 40 days prior to dispatch with a vaccine which covers all serotypes whose presence in the source population has been demonstrated through a surveillance programme as described in Appendix 3.8.X., and that the animals are identified in the accompanying certification as having been vaccinated; or
   b) the animals are not vaccinated, and a surveillance programme as described in Appendix X.X.X. has been in place in the source population for a period of at least 40 days immediately prior to dispatch, and no evidence of AHS has been detected.
4. An AHS free country or zone should be protected from an adjacent infected country or zone by a buffer zone in which surveillance is conducted as described in Appendix X.X.X.

Article 2.x.x.3.

AHS seasonally free zone

1. An AHS seasonally free zone is a part of an infected country or zone for which for part of a year, surveillance and monitoring demonstrate no evidence either of AHS transmission or of adult Culicoides likely to be competent AHS vectors.

2. For the application of Articles 2.x.x.7., 2.x.x. 10. and 2.x.x. 14., the seasonally free period is taken to commence the day following the last evidence of AHS transmission (as demonstrated by the surveillance programme), or of the cessation of activity of adult Culicoides likely to be competent AHS vectors.

3. For the application of Articles 2.x.x.7., 2.x.x. 10. and 2.x.x. 14., the seasonally free period is taken to conclude either:

   a) at least 28 days before the earliest date that historical data show AHS virus activity has recommenced; or

   b) immediately if current climatic data or data from a surveillance and monitoring programme indicate an earlier resurgence of activity of adult Culicoides likely to be competent AHS vectors.

4. An AHS seasonally free zone in which surveillance and monitoring has found no evidence that Culicoides likely to be competent AHS vectors are present will not lose its free status through the importation of vaccinated or seropositive animals, semen or embryos from infected countries or zones.

5. An AHS seasonally free zone in which surveillance and monitoring has found evidence that Culicoides likely to be competent AHS vectors are present will not lose its free status through the importation of vaccinated or seropositive domestic horses from infected countries or zones, provided:

   a) the animals have been vaccinated in accordance with the Terrestrial Manual at least 40 days prior to dispatch with a vaccine which covers all serotypes whose presence in the source population has been demonstrated through a surveillance programme as described in Appendix 3.8.X., and that the animals are identified in the accompanying certification as having been vaccinated; or

   b) the animals are not vaccinated, and a surveillance programme as described in Appendix X.X.X. has been in place in the source population for a period of at least 40 days immediately prior to dispatch, and no evidence of AHS has been detected.

Article 2.x.x.4.

AHS infected country or zone

An AHS infected country or zone is a clearly defined area where evidence of AHS has been reported during the past 2 years.
Appendix XXXI (contd)

Article 2.x.x.5.

_Veterinary Administrations_ of countries shall consider whether there is a risk with regard to AHS infection in accepting importation or transit through their territory, from other countries, of the following commodities:

1. equines;
2. equine semen;
3. equine embryos;
4. pathological material and biological products (from these species) (see Chapter 1.4.5. and Section 1.5.).

Other commodities should be considered as not having the potential to spread AHS when they are the subject of international trade.

Article 2.x.x.6.

When importing from AHS free countries or zones, _Veterinary Administrations_ should require:

_for domestic horses_

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 AHS free country or zone since birth or for at least 40 days prior to shipment;

**AND**

4. either:
   a) did not transit through an infected country or zone; or
   b) were protected from attack from _Culicoides_ likely to be competent AHS vectors at all times when transiting through an infected country or zone.

Article 2.x.x.7.

When importing from AHS free countries or zones, _Veterinary Administrations_ should require:

_for other equines_

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 AHS free country or zone since birth or for at least 40 days prior to shipment;
AND

if the animal originates from a zone or country adjacent to a zone or country considered infected with AHS:

4. were protected from attack from Culicoides likely to be competent AHS vectors for at least 40 days prior to shipment; and, either:
   a) were subjected during that period to a serological test according to the Terrestrial Manual to detect antibody to the AHS group, with negative results on two occasions, with an interval of not less than 7 days between each test, the first test being carried out at least 21 days after introduction into the quarantine station; or
   b) were subjected during that period to an agent identification test according to the Terrestrial Manual with negative results, on blood samples taken on two occasions, with an interval of not less than 7 days between each test, the first test being carried out at least 7 days after introduction into the quarantine station;

5. were protected from attack from Culicoides likely to be competent AHS vectors during transportation to and at the place of shipment.

Article 2.x.x.8.

When importing from AHS seasonally free zones, Veterinary Administrations should require:

for domestic horses

the presentation of an international veterinary certificate attesting that the animals:

1. were kept during the seasonally free period in an AHS seasonally free zone for at least 40 days prior to shipment;

2. have not been vaccinated against AHS within the past 40 days;

AND

3. either:
   a) did not transit through an infected country or zone; or
   b) were protected from attack from Culicoides likely to be competent AHS vectors at all times when transiting through an infected country or zone.

Article 2.x.x.9.

When importing from AHS infected countries or zones, Veterinary Administrations should require:

for domestic horses

the presentation of an international veterinary certificate attesting that the animals:
were protected from attack from *Culicoides* likely to be competent AHS vectors for at least 40 days prior to shipment; or

2. were protected from attack from *Culicoides* likely to be competent AHS vectors for at least 28 days prior to shipment, and were subjected during that period to a serological test in accordance with the *Terrestrial Manual* to detect antibody to the AHS group, with negative results on two occasions, with an interval of not less than 7 days between each test, the first test being carried out at least 21 days after introduction into the quarantine station; or

3. were protected from attack from *Culicoides* likely to be competent AHS vectors for at least 14 days prior to shipment, and were subjected during that period to an agent identification test in accordance with the *Terrestrial Manual* with negative results, on blood samples taken on two occasions, with an interval of not less than 7 days between each test, the first test being carried out at least 7 days after introduction into the quarantine station; AND

4. have not been vaccinated against AHS within the last 40 days;

5. were protected from attack from *Culicoides* likely to be competent AHS vectors during transportation to and at the place of shipment.

**Article 2.x.x.10.**

When importing from AHS free countries or zones, Veterinary Administrations should require:

for semen of domestic horses

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 within 40 days of the day of collection;
3. were kept in an AHS free country or zone for at least 40 days before commencement of, and during, collection of the semen.

**Article 2.x.x.11.**

When importing from AHS seasonally free zones, Veterinary Administrations should require:

for semen of domestic horses

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. were not vaccinated against AHS within 40 days of the day of collection;
3. were kept during the seasonally free period in an AHS seasonally free zone for at least 40 days before commencement of, and during, collection of the semen.
Article 2.x.x.12.

When importing from AHS infected countries or zones, Veterinary Administrations should require:

for semen of domestic horses

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. were not vaccinated against AHS within 40 days of the day of collection;
3. were protected from attack from Culicoides likely to be competent AHS vectors for at least 40 days before commencement of, and during, collection of the semen.

Article 2.x.x.13.

When importing from AHS free countries or zones, Veterinary Administrations should require:

for in vivo derived embryos of domestic horses

the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) showed no clinical sign of AHS on the day of collection of the embryos and for the following 40 days;
   b) have not been vaccinated against AHS within 40 days prior to collection;
   c) were kept in an AHS free country or zone for at least the 40 days prior to, and at the time of, embryo collection;
2. the embryos were collected, processed and stored in conformity with the provisions of Appendix 3.3.1.

Article 2.x.x.14.

When importing from AHS seasonally free zones, Veterinary Administrations should require:

for in vivo derived embryos of domestic horses

the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) showed no clinical sign of AHS on the day of collection of the embryos and for the following 40 days;
   b) have not been vaccinated against AHS within the 40 days prior to collection;
   c) were kept during the seasonally free period in an AHS seasonally free zone for at least the 40 days prior to, and at the time of, collection of the embryos;
2. the embryos were collected, processed and stored in conformity with the provisions of Appendix 3.3.1.

Article 2.x.x.15.

When importing from AHS infected countries or zones, Veterinary Administrations should require:

for in vivo derived embryos of domestic horses

the presentation of an international veterinary certificate attesting that:

1. the donor females:
   a) showed no clinical sign of AHS on the day of collection of the semen and for the following 40 days;
   b) have not been vaccinated against AHS within the 40 days prior to collection;
   c) were protected from attack from Culicoides likely to be competent AHS vectors for at least 40 days before commencement of, and during, collection of the embryos;

2. the embryos were collected, processed and stored in conformity with the provisions of Appendix 3.3.1.

Article 2.x.x.16.

Protecting animals from Culicoides attack

When transporting equines through AHS infected countries or zones, Veterinary Administrations should require strategies to protect animals from attack from Culicoides likely to be competent AHS vectors during transport, taking into account the local ecology of the vector.

Potential risk management strategies include:

1. treating animals with chemical repellents prior to and during transportation;
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 shadecloth;
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.
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