
of 29 April 2004

laying down specific rules for the organisation of official controls on products of animal origin intended for human consumption

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<table>
<thead>
<tr>
<th>No</th>
<th>page</th>
<th>date</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1</td>
<td>L 165</td>
<td>30.4.2004</td>
</tr>
<tr>
<td>M2</td>
<td>L 338</td>
<td>22.12.2005</td>
</tr>
<tr>
<td>M3</td>
<td>L 338</td>
<td>22.12.2005</td>
</tr>
<tr>
<td>M4</td>
<td>L 320</td>
<td>18.11.2006</td>
</tr>
<tr>
<td>M5</td>
<td>L 363</td>
<td>20.12.2006</td>
</tr>
<tr>
<td>M6</td>
<td>L 277</td>
<td>18.10.2008</td>
</tr>
<tr>
<td>M7</td>
<td>L 87</td>
<td>31.3.2009</td>
</tr>
<tr>
<td>M8</td>
<td>L 149</td>
<td>15.6.2010</td>
</tr>
<tr>
<td>M9</td>
<td>L 46</td>
<td>19.2.2011</td>
</tr>
<tr>
<td>M10</td>
<td>L 196</td>
<td>28.7.2011</td>
</tr>
<tr>
<td>M11</td>
<td>L 158</td>
<td>10.6.2013</td>
</tr>
<tr>
<td>M12</td>
<td>L 69</td>
<td>8.3.2014</td>
</tr>
<tr>
<td>M13</td>
<td>L 69</td>
<td>8.3.2014</td>
</tr>
<tr>
<td>M14</td>
<td>L 175</td>
<td>14.6.2014</td>
</tr>
<tr>
<td>M15</td>
<td>L 323</td>
<td>9.12.2015</td>
</tr>
<tr>
<td>M16</td>
<td>L 285</td>
<td>1.11.2017</td>
</tr>
</tbody>
</table>

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CHAPTER I
GENERAL PROVISIONS

Article 1
Scope

1. This Regulation lays down specific rules for the organisation of official controls on products of animal origin.

1a. This Regulation shall apply in addition to Regulation (EC) No 882/2004 of the European Parliament and of the Council of 29 April 2004 on official controls performed to ensure the verification of compliance with feed and food law, animal health and animal welfare rules (1).

2. It shall apply only in respect of activities and persons to which Regulation (EC) No 853/2004 applies.

3. The performance of official controls pursuant to this Regulation shall be without prejudice to food business operators' primary legal responsibility for ensuring food safety, as laid down in Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority, and laying down procedures in matters of food safety (2), and any civil or criminal liability arising from the breach of their obligations.

Article 2
Definitions

1. For the purposes of this Regulation, the following definitions shall apply:

(c) 'competent authority' means the central authority of a Member State competent to carry out veterinary checks or any authority to which it has delegated that competence;

(f) ‘official veterinarian’ means a veterinarian qualified, in accordance with this Regulation, to act in such a capacity and appointed by the competent authority;

(g) ‘approved veterinarian’ means a veterinarian designated by the competent authority to carry out specific official controls on holdings on its behalf;

(h) ‘official auxiliary’ means a person qualified, in accordance with this Regulation, to act in such a capacity, appointed by the competent authority and working under the authority and responsibility of an official veterinarian;

and

(i) ‘health mark’ means a mark indicating that, when it was applied, official controls had been carried out in accordance with this Regulation.

2. The definitions laid down in the following Regulations shall also apply as appropriate:

(a) Regulation (EC) No 178/2002;

(b) the definitions of ‘animal by-products’, ‘TSEs’ (transmissible spongiform encephalopathies) and ‘specified risk material’ laid down in Regulation (EC) No 1774/2002 of the European Parliament and of the Council of 3 October 2002 laying down health rules concerning animal by-products not intended for human consumption (1);

(b)(a) Regulation (EC) No 882/2004;

(c) Regulation (EC) No 852/2004, except for the definition of ‘competent authority’;

and


CHAPTER II
OFFICIAL CONTROLS IN RELATION TO COMMUNITY ESTABLISHMENTS

Article 3
Approval of establishments

1. The competent authorities shall approve establishments when, and in the manner, specified in Article 31(2) of Regulation (EC) No 882/2004.

2. In the case of factory and freezer vessels flying the flag of Member States, the maximum periods of three and six months applying to the conditional approval of other establishments may be extended, if necessary. However, conditional approval shall not exceed a total of 12 months. Inspections of such vessels shall take place as specified in Annex III.

3. The competent authority shall give each approved establishment, including those with conditional approval, an approval number, to which codes may be added to indicate the types of products of animal origin manufactured. For wholesale markets, secondary numbers indicating units or groups of units selling or manufacturing products of animal origin may be added to the approval number.

4. (c) In the case of wholesale markets, the competent authority may withdraw or suspend approval in respect of certain units or groups of units.

5. Paragraphs 1, 2 and 3 shall apply both:

(a) to establishments that begin placing products of animal origin on the market on or after the date of application of this Regulation; and

(b) to establishments already placing products of animal origin on the market but in respect of which there was previously no requirement for approval. In the latter case, the competent authority’s on-site visit required under paragraph 1 shall take place as soon as possible.

Paragraph 4 shall also apply to approved establishments that placed products of animal origin on the market in accordance with Community legislation immediately prior to the application of this Regulation.

Article 4

General principles for official controls in respect of all products of animal origin falling within the scope of this Regulation

1. Member States shall ensure that food business operators offer all assistance needed to ensure that official controls carried out by the competent authority can be performed effectively.

They shall in particular:

— give access to all buildings, premises, installations or other infrastructures;

— make available any documentation and record required under the present regulation or considered necessary by the competent authority for judging the situation.

2. The competent authority shall carry out official controls to verify food business operators’ compliance with the requirements of:

(a) Regulation (EC) No 852/2004;

(b) Regulation (EC) No 853/2004;

and

(c) Regulation (EC) No 1774/2002.
3. The official controls referred to in paragraph 1 shall include:

(a) audits of good hygiene practices and hazard analysis and critical control point (HACCP)-based procedures;

(b) the official controls specified in Articles 5 to 8;

and

(c) any particular auditing tasks specified in the Annexes.

4. Audits of good hygiene practices shall verify that food business operators apply procedures continuously and properly concerning at least:

(a) checks on food-chain information;

(b) the design and maintenance of premises and equipment;

(c) pre-operational, operational and post-operational hygiene;

(d) personal hygiene;

(e) training in hygiene and in work procedures;

(f) pest control;

(g) water quality;

(h) temperature control;

and

(i) controls on food entering and leaving the establishment and any accompanying documentation.

5. Audits of HACCP-based procedures shall verify that food business operators apply such procedures continuously and properly, having particular regard to ensuring that the procedures provide the guarantees specified in Section II of Annex II to Regulation (EC) No 853/2004. They shall, in particular, determine whether the procedures guarantee, to the extent possible, that products of animal origin:

(a) comply with microbiological criteria laid down under Community legislation;

(b) comply with Community legislation on residues, contaminants and prohibited substances;

and

(c) do not contain physical hazards, such as foreign bodies.

When, in accordance with Article 5 of Regulation (EC) No 852/2004, a food business operator uses procedures set out in guides to the application of HACCP principles rather than establishing its own specific procedures, the audit shall cover the correct use of these guides.

6. Verification of compliance with the requirements of Regulation (EC) No 853/2004 concerning the application of identification marks shall take place in all establishments approved in accordance with that Regulation, in addition to verification of compliance with other traceability requirements.

7. In the case of slaughterhouses, game handling establishments and cutting plants placing fresh meat on the market, an official veterinarian shall carry out the auditing tasks referred to in paragraphs 3 and 4.
8. When carrying out auditing tasks, the competent authority shall take special care:

(a) to determine whether staff and staff activities in the establishment at all stages of the production process comply with the relevant requirements of the Regulations referred to in paragraph 1(a) and (b). To support the audit, the competent authority may carry out performance tests, in order to ascertain that staff performance meets specified parameters;

(b) to verify the food business operator's relevant records;

(c) to take samples for laboratory analysis whenever necessary;

and

(d) to document elements taken into account and the findings of the audit.

9. The nature and intensity of auditing tasks in respect of individual establishments shall depend upon the assessed risk. To this end, the competent authority shall regularly assess:

(a) public and, where appropriate, animal health risks;

(b) in the case of slaughterhouses, animal welfare aspects;

(c) the type and throughput of the processes carried out;

and

(d) the food business operator's past record as regards compliance with food law.

Article 5

Fresh meat

Member States shall ensure that official controls with respect to fresh meat take place in accordance with Annex I.

1. The official veterinarian shall carry out inspection tasks in slaughterhouses, game handling establishments and cutting plants placing fresh meat on the market in accordance with the general requirements of Section I, Chapter II, of Annex I, and with the specific requirements of Section IV, in particular as regards:

(a) food chain information;

(b) ante-mortem inspection;

(c) animal welfare;

(d) post-mortem inspection;

(e) specified risk material and other animal by-products;

and

(f) laboratory testing.
2. The health marking of carcases of domestic ungulates, farmed game mammals other than lagomorphs, and large wild game, as well as half-carcases, quarters and cuts produced by cutting half-carcases into three wholesale cuts, shall be carried out in slaughterhouses and game-handling establishments in accordance with Section I, Chapter III, of Annex I. Health marks shall be applied by, or under the responsibility of, the official veterinarian when official controls have not identified any deficiencies that would make the meat unfit for human consumption.

3. After carrying out the controls mentioned in points 1 and 2, the official veterinarian shall take appropriate measures as set out in Annex I, Section II, in particular as regards:

(a) the communication of inspection results;

(b) decisions concerning food chain information;

(c) decisions concerning live animals;

(d) decisions concerning animal welfare;

and

(e) decisions concerning meat.

4. Official auxiliaries may assist the official veterinarian with official controls carried out in accordance with Sections I and II of Annex I as specified in Section III, Chapter I. In that case, they shall work as part of an independent team.

5. (a) Member States shall ensure that they have sufficient official staff to carry out the official controls required under Annex I with the frequency specified in Section III, Chapter II.

(b) A risk-based approach shall be followed to assess the number of official staff that need to be present on the slaughter line in any given slaughterhouse. The number of official staff involved shall be decided by the competent authority and shall be such that all the requirements of this Regulation can be met.

6. (a) Member States may allow slaughterhouse staff to assist with official controls by carrying out certain specific tasks, under the supervision of the official veterinarian, in relation to the production of meat from poultry and lagomorphs in accordance with Annex I, Section III, Chapter III, part A. If they do so, they shall ensure that staff carrying out such tasks:

(i) are qualified and undergo training in accordance with those provisions;

(ii) act independently from production staff;

and

(iii) report any deficiency to the official veterinarian.

(b) Member States may also allow slaughterhouse staff to carry out specific sampling and testing tasks in accordance with Annex I, Section III, Chapter III, Part B.
7. Member States shall ensure that official veterinarians and official auxiliaries are qualified and undergo training in accordance with Annex I, Section III, Chapter IV.

Article 6

Live bivalve molluscs

Member States shall ensure that the production and placing on the market of live bivalve molluscs, live echinoderms, live tunicates and live marine gastropods undergo official controls as described in Annex II.

Article 7

Fishery products

Member States shall ensure that official controls with respect to fishery products take place in accordance with Annex III.

Article 8

Raw milk and dairy products

Member States shall ensure that official controls with respect to raw milk and dairy products take place in accordance with Annex IV.

CHAPTER III

PROCEDURES CONCERNING IMPORTS

Article 10

To ensure the uniform application of the principles and conditions laid down in Article 11 of Regulation (EC) No 178/2002 and Title VI, Chapter II, of Regulation (EC) No 882/2004 the procedures laid down in this Chapter shall apply.

Article 11

Lists of third countries and parts of third countries from which imports of specified products of animal origin are permitted

1. Products of animal origin shall be imported only from a third country or a part of third country that appears on a list drawn up and updated in accordance with the procedure referred to in Article 19(2).

2. A third country shall appear on such lists only if a Community control in that country has taken place and demonstrates that the competent authority provides appropriate guarantees as specified in Article 48(3) of Regulation (EC) No 882/2004. However, a third country may appear on such lists without a Community control having taken place if:
(a) the risk determined in accordance with Article 46(3)(a) of Regulation (EC) No 882/2004 does not warrant it;

and

(b) it is determined, when deciding to add a particular third country to a list in accordance with paragraph 1, that other information indicates that the competent authority provides the necessary guarantees.

3. Lists drawn up in accordance with this Article may be combined with other lists drawn up for public and animal health purposes.

4. When drawing up or updating lists, particular account shall be taken of the criteria listed in Articles 46 and 48(3) of Regulation (EC) No 882/2004. Regard shall also be had to:

(a) the legislation of the third country on:

(i) products of animal origin,

(ii) the use of veterinary medicinal products, including rules on their prohibition or authorisation, their distribution, their placing on the market and the rules covering administration and inspection;

and

(iii) the preparation and use of feedingstuffs, including the procedures for using additives and the preparation and use of medicated feedingstuffs, as well as the hygiene quality of the raw materials used for preparing feedingstuffs and of the final product;

(i) the hygiene conditions of production, manufacture, handling, storage and dispatch actually applied to products of animal origin destined for the Community;

(j) any experience of marketing of the product from the third country and the results of any import controls carried out;

(k) the results of Community controls carried out in the third country, in particular the results of the assessment of the competent authorities, and the action that competent authorities have taken in the light of any recommendations addressed to them following a Community control;

(l) the existence, implementation and communication of an approved zoonoses control programme;

and

(m) the existence, implementation and communication of an approved residue control programme.
5. The Commission shall arrange for up-to-date versions of all lists drawn up or updated in accordance with this Article to be available to the public.

Article 12
List of establishments from which imports of specified products of animal origin are permitted

1. Products of animal origin may be imported into the Community only if they have been dispatched from, and obtained or prepared in, establishments that appear on lists drawn up and updated in accordance with this Article, except:

(a) when, on a case-by-case basis, it is decided, in accordance with the procedure referred to in Article 19(2), that the guarantees that a specified third country provides in respect of imports of specified products of animal origin are such that the procedure provided for in this Article is unnecessary to ensure compliance with the requirements of paragraph 2;

and

(b) in the cases specified in Annex V.

In addition, fresh meat, minced meat, meat preparations, meat products and mechanically separated meat (MSM) may be imported into the Community only if they have been manufactured from meat obtained in slaughterhouses and cutting plants appearing on lists drawn up and updated in accordance with this Article or in approved Community establishments.

2. An establishment may be placed on such a list only if the competent authority of the third country of origin guarantees that:

(a) that establishment, together with any establishments handling raw material of animal origin used in the manufacture of the products of animal origin concerned, complies with relevant Community requirements, in particular those of Regulation (EC) No 853/2004, or with requirements that were determined to be equivalent to such requirements when deciding to add that third country to the relevant list in accordance with Article 11;

(b) an official inspection service in that third country supervises the establishments and makes available to the Commission, where necessary, all relevant information on establishments furnishing raw materials;

and

(c) it has real powers to stop the establishments from exporting to the Community in the event that the establishments fail to meet the requirements referred to under (a).

3. The competent authorities of third countries appearing on lists drawn up and updated in accordance with Article 11 shall guarantee that lists of the establishments referred to in paragraph 1 are drawn up, kept up-to-date and communicated to the Commission.
4. (a) The Commission shall provide the contact points that Member States have designated for this purpose with regular notifications concerning new or updated lists that it has received from the competent authorities of third countries concerned in accordance with paragraph 3.

(b) If no Member State objects to the new or updated list within 20 working days of the Commission's notification, imports shall be authorised from establishments appearing on the list 10 working days after the day on which the Commission makes it available to the public.

(c) The Commission shall, whenever at least one Member State makes written comments, or whenever it considers that the modification of a list is necessary in the light of relevant information such as Community inspection reports or a notification under the rapid alert system, inform all Member States and include the point on agenda of the next meeting of the relevant section of the Standing Committee on the Food Chain and Animal Health for decision, where appropriate, in accordance with the procedure referred to in Article 19(2).

5. The Commission shall arrange for up-to-date versions of all lists to be available to the public.

**Article 13**

**Live bivalve molluscs, echinoderms, tunicates and marine gastropods**

1. Notwithstanding Article 12(1)(b), live bivalve molluscs, echinoderms, tunicates and marine gastropods shall come from production areas in third countries that appear on lists drawn up and updated in accordance with Article 12.

2. The requirement of paragraph 1 shall not apply to pectinidae harvested outside classified production areas. However, official controls with respect to pectinidae shall take place in accordance with Annex II, Chapter III.

3. (a) Before the lists referred to in paragraph 1 are drawn up, particular account shall be taken of the guarantees that the competent authority of the third country can give concerning compliance with the requirements of this Regulation on the classification and control of production zones.

(b) An on-the-spot Community inspection visit shall take place before such lists are drawn up unless:

   (i) the risk determined in accordance with Article 18(18) does not warrant it;

and

   (ii) it is determined, when deciding to add a particular production area to a list in accordance with paragraph 1, that other information indicates that the competent authority provides the necessary guarantees.
4. The Commission shall arrange for up-to-date versions of all lists drawn up or updated in accordance with this Article to be available to the public.

Article 14

Documents

1. A document meeting the requirements set out in Annex VI shall accompany consignments of products of animal origin when they are imported into the Community.

2. The document shall certify that the products satisfy:

(a) the requirements laid down for such products according to Regulation (EC) No 852/2004 and Regulation (EC) No 853/2004 or provisions that are equivalent to those requirements;

and

(b) any specific import conditions established in accordance with Article 48 of Regulation (EC) No 882/2004.

3. Documents may include details required in accordance with other Community legislation on public and animal health matters.

4. Exemptions from paragraph 1 may be granted in accordance with the procedure referred to in Article 19(2) when it is possible to obtain the guarantees referred to in paragraph 2 of this Article in another manner.

Article 15

Special provisions for fishery products

1. The procedures laid down in this Chapter do not apply to fresh fishery products landed in the Community directly from a fishing vessel flying the flag of a third country.

Official controls with respect to such fishery products shall take place in accordance with Annex III.

2. (a) Fishery products imported from a factory or freezer vessel flying the flag of a third country shall come from vessels that appear on a list drawn up and updated in accordance with the procedure set out in Article 12(4).

(b) However, by way of exemption from Article 12(2)(b), a vessel may also be included on such lists:

(i) on the basis of a joint communication from the competent authority of the third country the flag of which the vessel is flying and from the competent authority of another third country to which the former competent authority has delegated responsibility for the inspection of the vessel concerned, on condition that:

— that third country appears on the list of third countries, drawn up in accordance with Article 11, from which imports of fisheries products are permitted,
— all fishery products from the vessel concerned that are
destined for placing on the market in the Community
are landed directly in that third country,
— the competent authority of that third country has
inspected the vessel and has declared that it complies
with Community requirements,
and
— the competent authority of that third country has
declared that it will regularly inspect the vessel to
ensure that it continues to comply with Community
requirements;
or
(ii) on the basis of a joint communication from the competent
authority of the third country the flag of which the vessel
is flying and from the competent authority of a Member
State, to which the former competent authority has
delegated responsibility for the inspection of the vessel
concerned, on condition that:
— all fishery products from the vessel concerned that are
destined for placing on the market in the Community
are landed directly in that Member State,
— the competent authority of that Member State has
inspected the vessel and has declared that it complies
with Community requirements,
and
— the competent authority of that Member State has
declared that it will regularly inspect the vessel to
ensure that it continues to comply with Community
requirements.

(c) The Commission shall arrange for up-to-date versions of all
lists drawn up or updated in accordance with this Article to be
available to the public.

3. When fishery products are imported directly from a fishing or
freezer vessel, a document signed by the captain may replace the
document required under Article 14.

4. Detailed rules for the implementation of this Article may be laid
down in accordance with the procedure referred to in Article 19(2).

CHAPTER IV

FINAL PROVISIONS

Article 16

Transitional measures of general scope designed to amend non-essential
elements of this Regulation, inter alia, by supplementing it with new
non-essential elements, in particular further specifications of the
requirements laid down in this Regulation, shall be adopted in
accordance with the regulatory procedure with scrutiny referred to in
Article 19(3).
Other implementing or transitional measures may be adopted in accordance with the regulatory procedure referred to in Article 19(2).

**Article 17**

Amendment and adaptation of the Annexes

1. Annexes I, II, III, IV, V and VI may be amended or supplemented by the Commission to take account of scientific and technical progress. Those measures, designed to amend non-essential elements of this Regulation, *inter alia*, by supplementing it, shall be adopted in accordance with the regulatory procedure with scrutiny referred to in Article 19(3).

2. Exemptions from Annexes I, II, III, IV, V and VI may be granted by the Commission, provided that they do not affect the achievement of the objectives of this Regulation. Those measures, designed to amend non-essential elements of this Regulation by supplementing it, shall be adopted in accordance with the regulatory procedure with scrutiny referred to in Article 19(3).

3. Member States may, without compromising achievement of the objectives of this Regulation, adopt, in accordance with paragraphs 4 to 7, national measures adapting the requirements laid down in Annex I.

4. The national measures referred to in paragraph 3 shall:

   (a) have the aim of:

      (i) enabling the continued use of traditional methods at any of the stages of production, processing or distribution of food;

      (ii) accommodating the needs of food businesses with a low throughput or that are situated in regions that are subject to special geographic constraints;

   or

      (iii) permitting pilot projects to take place in order to try out new approaches to hygiene controls on meat;

   (b) concern in particular the following elements of Annex I:

      (i) food chain information;

      (ii) the presence of the competent authority in establishments.

5. Any Member State wishing to adopt national measures as referred to in paragraph 3 shall notify the Commission and other Member States. Each notification shall:

   (a) provide a detailed description of the requirements that that Member State considers need to be adapted and the nature of the adaptation sought;

   (b) describe the establishments concerned;
(c) explain the reasons for the adaptation, including, where relevant, by providing a summary of the hazard analysis carried out and any measures to be taken to ensure that the adaptation will not compromise the objectives of this Regulation;

and

(d) give any other relevant information.

6. The other Member States shall have three months from the receipt of a notification referred to in paragraph 5 to send written comments to the Commission. The Commission may, and when it receives written comments from one or more Member States shall, consult Member States within the committee referred to in Article 19(1). The Commission may decide, in accordance with the procedure referred to in Article 19(2), whether the envisaged measures may be implemented subject, if necessary, to appropriate amendments. Where appropriate, the Commission may propose general measures in accordance with paragraphs 1 or 2 of this Article.

7. A Member State may adopt national measures adapting the requirements of Annex I only:

(a) in compliance with a decision adopted in accordance with paragraph 6;

(b) if, one month after the expiry of the period referred to in paragraph 6, the Commission has not informed Member States that it has received written comments or that it intends to propose the adoption of a decision in accordance with paragraph 6.

8. When a Member State adopts national measures implementing a pilot project to try out new approaches to hygiene controls on meat in accordance with paragraphs 3 to 7, the Member State shall communicate the results to the Commission as soon as they are available. The Commission shall then consider proposing general measures in accordance with paragraph 1.

**Article 18**

**Specific decisions**

Without prejudice to the general application of Article 16 and Article 17(1), implementing measures may be laid down in accordance with the regulatory procedure referred to in Article 19(2), and amendments to Annexes I, II, III, IV, V or VI, as measures designed to amend non-essential elements of this Regulation, may be adopted in accordance with the regulatory procedure with scrutiny referred to in Article 19(3), to specify:

1. tests to assess the performance of food business operators and their staff;

2. the method of communicating inspection results;

3. criteria to determine when, on the basis of a risk analysis, the official veterinarian need not be present in slaughterhouses and game handling establishments throughout ante-mortem and post-mortem inspection;
4. rules concerning the content of tests for official veterinarians and official auxiliaries;

5. microbiological criteria for process control in relation to hygiene in establishments;

6. alternative procedures, serological or other laboratory tests that provide guarantees at least equivalent to specific post-mortem inspection procedures described in Annex I, Section IV, and may therefore replace them, if the competent authority so decides;

7. circumstances in which certain of the specific post-mortem inspection procedures described in Annex I, Section IV, are not necessary, having regard to the holding, region or country of origin and to the principles of risk analysis,

8. rules for laboratory testing;

9. the cold treatment to be applied to meat in relation to cysticercosis and trichinosis;

10. conditions under which holdings and regions can be certified as officially free of cysticercus or trichinae;

11. methods to be applied when examining for the conditions referred to in Annex I, Section IV, Chapter IX;

12. for fattening pigs, criteria for controlled housing conditions and integrated production systems;

13. criteria for the classification of production and relaying areas for live bivalve molluscs in cooperation with the relevant Community Reference Laboratory, including:

   (a) limit values and analysis methods for marine biotoxins,

   (b) virus testing procedures and virological standards,

   and

   (c) sampling plans and the methods and analytical tolerances to be applied to check compliance with the criteria;

14. organoleptic criteria for the evaluation of the freshness of fishery products;

15. analytical limits, methods of analysis and sampling plans for the official controls on fishery products required under Annex III, including with regard to parasites and environmental contaminants;

16. the method by which the Commission will make lists of third countries and establishments in third countries available to the public pursuant to Articles 11, 12, 13 and 15.
Article 19

Committee procedure


2. Where reference is made to this paragraph, Articles 5 and 7 of Decision 1999/468/EC shall apply, having regard to the provisions of Article 8 thereof.

The period laid down in Article 5(6) of Decision 1999/468/EC shall be set at three months.

3. Where reference is made to this paragraph, Article 5a(1) to (4) and Article 7 of Decision 1999/468/EC shall apply, having regard to the provisions of Article 8 thereof.

Article 20

Consultation of the European Food Safety Authority

The Commission shall consult the European Food Safety Authority on matters falling within the scope of this Regulation whenever necessary and, in particular:

1. before proposing to modify the specific requirements concerning post-mortem inspection procedures laid down in Section IV of Annex I;

2. before proposing to modify the rules of Annex I, Section IV, Chapter IX, on meat from animals in which post-mortem inspection has revealed lesions indicating infection with brucellosis or tuberculosis; and

3. before proposing implementing measures on the matters referred to in Article 18(5) to (15).

Article 21

Report to the European Parliament and to the Council

1. The Commission shall, not later than 20 May 2009, submit a report to the European Parliament and the Council reviewing the experience gained from the application of this Regulation.

2. The Commission shall, if appropriate, accompany the report with relevant proposals.

Article 22

Entry into force

This Regulation shall enter into force on the 20th day after that of its publication in the Official Journal of the European Union.

It shall apply 18 months after the date on which all of the following acts have entered into force:
(a) Regulation (EC) No 852/2004;
(b) Regulation (EC) No 853/2004
and

However, it shall apply no earlier than 1 January 2006.

This Regulation shall be binding in its entirety and directly applicable in all Member States.

ANNEX I

FRESH MEAT

SECTION I: TASKS OF THE OFFICIAL VETERINARIAN

CHAPTER I: AUDITING TASKS

1. In addition to the general requirements of Article 4(4) concerning audits of good hygiene practices, the official veterinarian is to verify continuous compliance with food business operators' own procedures concerning any collection, transport, storage, handling, processing and use or disposal of animal by-products, including specified risk material, for which the food business operator is responsible.

2. In addition to the general requirements of Article 4(5) concerning audits of HACCP-based principles, the official veterinarian is to check that the operators' procedures guarantee, to the extent possible, that meat:

   (a) does not contain patho-physiological abnormalities or changes;

   (b) does not bear faecal or other contamination;

   and

   (c) does not contain specified risk material, except as provided for under Community legislation, and has been produced in accordance with Community legislation on TSEs.

CHAPTER II: INSPECTION TASKS

When carrying out inspection tasks in accordance with this Chapter, the official veterinarian is to take account of the results of the auditing tasks carried out in accordance with Article 4 and Chapter I of this Annex. Where appropriate he or she is to target inspection tasks accordingly.

A. Food chain information

1. The official veterinarian is to check and analyse relevant information from the records of the holding of provenance of animals intended for slaughter and to take account of the documented results of this check and analysis when carrying out ante- and post-mortem inspection.

2. When carrying out inspection tasks, the official veterinarian is to take account of official certificates accompanying animals, and any declarations made by veterinarians carrying out controls at the level of primary production, including official veterinarians and approved veterinarians.

3. When food business operators in the food chain take additional measures to guarantee food safety by implementing integrated systems, private control systems, independent third party certification or by other means, and when these measures are documented and animals covered by these schemes clearly identifiable, the official veterinarian may take this into account when carrying out inspection tasks and reviewing the HACCP-based procedures.

B. Ante-mortem inspection

1. Subject to paragraphs 4 and 5:

   (a) the official veterinarian is to carry out an ante-mortem inspection of all animals before slaughter;

   (b) that inspection must take place within 24 hours of arrival at the slaughterhouse and less than 24 hours before slaughter.

In addition, the official veterinarian may require inspection at any other time.
2. Ante-mortem inspection must in particular determine whether, as regards the particular animal inspected, there is any sign:

(a) that welfare has been compromised;

or

(b) of any condition which might adversely affect human or animal health, paying particular attention to the detection of zoonotic diseases and animal diseases for which animal health rules are laid down in Union legislation.

3. In addition to routine ante-mortem inspection, the official veterinarian is to carry out a clinical inspection of all animals that the food business operator or an official auxiliary may have put aside.

4. In the case of emergency slaughter outside the slaughterhouse and of hunted wild game, the official veterinarian at the slaughterhouse or game handling establishment is to examine the declaration accompanying the body of the animal issued by the veterinarian or the trained person in accordance with Regulation (EC) No 853/2004.

5. Where provided for in Section III, Chapter II, or in Section IV, ante-mortem inspection may be carried out at the holding of provenance. In such cases, the official veterinarian at the slaughterhouse need carry out ante-mortem inspection only when and to the extent specified.

C. Animal welfare

The official veterinarian is to verify compliance with relevant Community and national rules on animal welfare, such as rules concerning the protection of animals at the time of slaughter and during transport.

D. Post-mortem inspection

1. Carcases and accompanying offal must be subjected without delay after slaughter to post-mortem inspection. All external surfaces must be viewed. Minimal handling of the carcases and offal or special technical facilities may be required for that purpose. Particular attention must be paid to the detection of zoonotic diseases and animal diseases for which animal health rules are laid down in Union legislation. The speed of the slaughter line and the number of inspection staff present must be such as to allow for proper inspection.

2. Additional examinations are to take place, such as palpation and incision of parts of the carcase and offal and laboratory tests, whenever considered necessary:

(a) to reach a definitive diagnosis;

or

(b) to detect the presence of:

(i) an animal disease,

(ii) residues or contaminants in excess of the levels laid down under Community legislation,

(iii) non-compliance with microbiological criteria,

or

(iv) other factors that might require the meat to be declared unfit for human consumption or restrictions to be placed on its use, particularly in the case of animals having undergone emergency slaughter.
3. The official veterinarian is to require carcases of domestic solipeds, bovine animals over six months old, and domestic swine over four weeks old to be submitted for post-mortem inspection split lengthways into half carcases down the spinal column. If the inspection so necessitates, the official veterinarian may also require any head or any carcase to be split lengthways. However, to take account of particular eating habits, technological developments or specific sanitary situations, the competent authority may authorise the submission for inspection of carcases of domestic solipeds, bovine animals over six months old, and domestic swine over four weeks old, not split in half.

4. During the inspection, precautions must be taken to ensure that contamination of the meat by actions such as palpation, cutting or incision is kept to a minimum.

5. In the event of an emergency slaughter, the carcase shall be subjected to post-mortem examination as soon as possible in accordance with paragraphs 1 to 4 before it is released for human consumption.

E. Specified risk material and other animal by-products

In accordance with specific Community rules on specified risk material and other animal by-products, the official veterinarian is to check the removal, separation and, where appropriate, marking of such products. The official veterinarian is to ensure that the food business operator takes all necessary measures to avoid contaminating meat with specified risk material during slaughter (including stunning) and removal of specified risk material.

F. Laboratory testing

1. The official veterinarian is to ensure that sampling takes place and that samples are appropriately identified and handled and sent to the appropriate laboratory within the framework of:

   (a) the monitoring and control of zoonoses and zoonotic agents;

   (b) specific laboratory testing for the diagnosis of TSEs in accordance with Regulation (EC) No 999/2001 of the European Parliament and of the Council (1);

   (c) the detection of unauthorised substances or products and the control of regulated substances, in particular within the framework of the National Residue Plans referred to in Council Directive 96/23/EC (2);

   and

   (d) the detection of animal diseases for which animal health rules are laid down in Union legislation.

2. The official veterinarian is also to ensure that any other necessary laboratory testing takes place.

CHAPTER III: HEALTH MARKING

1. The official veterinarian is to supervise health marking and the marks used.

2. The official veterinarian is to ensure, in particular, that:

(a) the health mark is applied only to animals (domestic ungulates, farmed game mammals other than lagomorphs, and large wild game) having undergone ante-mortem and post-mortem inspection in accordance with this Regulation and when there are no grounds for declaring the meat unfit for human consumption. However, the health mark may be applied before the results of any examination for trichinosis is available, if the official veterinarian is satisfied that meat from the animal concerned will be placed on the market only if the results are satisfactory;

and

(b) health-marking takes place on the external surface of the carcase, by stamping the mark in ink or hot branding, and in such a manner that, if carcases are cut into half carcases or quarters, or half carcases are cut into three pieces, each piece bears a health mark.

3. The health mark must be an oval mark at least 6,5 cm wide by 4,5 cm high bearing the following information in perfectly legible characters:

(a) the mark must indicate name of the country in which the establishment is located, which may be written out in full in capitals or shown as a two-letter code in accordance with the relevant ISO standard.

In the case of Member States, however, these codes are BE, BG, CZ, DK, DE, EE, GR, ES, FR, HR, IE, IT, CY, LV, LT, LU, HU, MT, NL, AT, PL, PT, SI, SK, FI, RO, SE and UK.

(b) the mark must indicate the approval number of the slaughterhouse;

and

(c) When applied in an establishment located within the Community, the mark must be oval in shape and include the abbreviation CE, EC, EF, EG, EK, EO, EV, ES, EÜ, EK, EB, EZ or WE.

Those abbreviations must not be included in marks applied on meat imported into the Community from slaughterhouses located outside the Community.

4. Letters must be at least 0,8 cm high and figures at least 1 cm high. The dimensions and characters of the mark may be reduced for health marking of lamb, kids and piglets.

5. The colours used for health marking must be authorised in accordance with Community rules on the use of colouring substances in foodstuffs.

6. The health mark may also include an indication of the official veterinarian who carried out the health inspection of the meat.

8. Meat from unskinned wild game cannot bear a health mark unless, after skinning in a game handling establishment, it has undergone post-mortem inspection and been declared fit for human consumption.

9. This Chapter is to apply without prejudice to animal health rules on health marking.
SECTION II: ACTION FOLLOWING CONTROLS

CHAPTER I: COMMUNICATION OF INSPECTION RESULTS

1. The official veterinarian is to record and to evaluate the results of inspection activities.

2. (a) If inspections reveal the presence of any disease or condition that might affect public or animal health, or compromise animal welfare, the official veterinarian is to inform the food business operator.

   (b) When the problem identified arose during primary production, the official veterinarian attending the holding of provenance, the food business operator responsible for the holding of provenance (provided that such information would not prejudice subsequent legal proceedings) and, where appropriate, the competent authority responsible for supervising the holding of provenance or the hunting area.

   (c) If the animals concerned were raised in another Member State or in a third country, the official veterinarian is to inform the competent authority of the Member State where the establishment is located. That competent authority is to take appropriate measures in accordance with applicable Community legislation.

3. The results of inspections and tests are to be included in relevant databases.

4. When the official veterinarian, while carrying out ante-mortem or post-mortem inspection or any other inspection activity, suspects the presence of an infectious agent of animal diseases for which animal health rules are laid down in Union legislation, the official veterinarian must notify as appropriate the competent authority and both must take all necessary measures and precautions to prevent the possible spread of the infectious agent in accordance with applicable Union legislation.

CHAPTER II: DECISIONS CONCERNING FOOD CHAIN INFORMATION

1. The official veterinarian is to verify that animals are not slaughtered unless the slaughterhouse operator has been provided with and checked relevant food chain information.

2. However, the official veterinarian may allow animals to undergo slaughter in the slaughterhouse even if the relevant food chain information is not available. In this case, all relevant food chain information must be supplied before the carcase is approved for human consumption. Pending a final judgement, such carcases and related offal must be stored separately from other meat.

3. Notwithstanding paragraph 2, when relevant food chain information is not available within 24 hours of an animal’s arrival at the slaughterhouse, all meat from the animal is to be declared unfit for human consumption. If the animal has not yet been slaughtered, it is to be killed separately from other animals.

4. When the accompanying records, documentation or other information shows that:

   (a) animals come from a holding or an area subject to a movement prohibition or other restriction for reasons of animal or public health;

   (b) rules on the use of veterinary medicinal products have not been complied with;
(c) any other condition which might adversely affect human or animal health is present, animals may not be accepted for slaughter other than in accordance with procedures laid down under Community legislation to eliminate human or animal health risks.

If the animals are already present at the slaughterhouse, they must be killed separately and declared unfit for human consumption, taking precautions to safeguard animal and public health where appropriate. Whenever the official veterinarian considers it necessary, official controls are to be carried out on the holding of provenance.

5. The competent authority is to take appropriate action if it discovers that the accompanying records, documentation or other information do not correspond with the true situation on the holding of provenance or the true condition of the animals or aim deliberately to mislead the official veterinarian. The competent authority is to take action against the food business operator responsible for the holding of provenance of the animals, or any other person involved. This action may consist in particular of extra controls. The food business operator responsible for the holding of provenance or any other person involved are to bear the costs of such extra controls.

CHAPTER III: DECISIONS CONCERNING LIVE ANIMALS

1. The official veterinarian is to verify compliance with the food business operator's duty pursuant to Regulation (EC) No 853/2004 to ensure that animals accepted for slaughter for human consumption are properly identified. The official veterinarian is to ensure that animals whose identity is not reasonably ascertainable are killed separately and declared unfit for human consumption. Whenever the official veterinarian considers it necessary, official controls are to be carried out on the holding of provenance.

2. When there are overriding animal welfare considerations, horses may undergo slaughter at the slaughterhouse even if the legally required information concerning their identity has not been supplied. However, this information must be supplied before the carcase may be declared fit for human consumption. These requirements also apply in the case of emergency slaughter of horses outside the slaughterhouse.

3. The official veterinarian is to verify compliance with the food business operator's duty under Regulation (EC) No 853/2004 to ensure that animals that have such hide, skin or fleece conditions that there is an unacceptable risk of contamination of the meat during slaughter are not slaughtered for human consumption unless they are cleaned beforehand.

4. Animals with a disease or condition that may be transmitted to animals or humans through handling or eating meat and, in general, animals showing clinical signs of systemic disease or emaciation, are not to be slaughtered for human consumption. Such animals must be killed separately, under conditions such that other animals or carcases can not be contaminated, and declared unfit for human consumption.

5. The slaughter of animals suspected of having a disease or condition that may adversely affect human or animal health is to be deferred. Such animals are to undergo detailed ante-mortem examination in order to make a diagnosis. In addition, the official veterinarian may decide that sampling and laboratory examinations are to take place to supplement post-mortem inspection. If necessary, the animals are to be slaughtered separately or at the end of normal slaughtering, taking all necessary precautions to avoid contamination of other meat.
6. Animals that might contain residues of veterinary medicinal products in excess of the levels laid down in accordance with Community legislation, or residues of forbidden substances, are to be dealt with in accordance with Directive 96/23/EC.

7. The official veterinarian is to impose the conditions under which animals are to be dealt with under a specific scheme for the eradication or control of a specific disease, such as brucellosis or tuberculosis, or zoonotic agents such as salmonella, under his/her direct supervision. The competent authority is to determine the conditions under which such animals may be slaughtered. These conditions must have the aim of minimising contamination of other animals and the meat of other animals.

8. Animals that are presented to a slaughterhouse for slaughter must as a general rule be slaughtered there. However, in exceptional circumstances, such as a serious breakdown of the slaughter facilities, the official veterinarian may allow direct movements to another slaughterhouse.

CHAPTER IV: DECISIONS CONCERNING ANIMAL WELFARE

1. When the rules concerning the protection of animals at the time of slaughter or killing are not respected, the official veterinarian is to verify that the food business operator immediately takes necessary corrective measures and prevents recurrence.

2. The official veterinarian is to take a proportionate and progressive approach to enforcement action, ranging from issuing directions to slowing down and stopping production, depending on the nature and gravity of the problem.

3. Where appropriate, the official veterinarian is to inform other competent authorities of welfare problems.

4. When the official veterinarian discovers that rules concerning the protection of animals during transport are not being respected, he or she is to take necessary measures in accordance with the relevant Community legislation.

5. When:

   (a) an official auxiliary is carrying out checks on animal welfare pursuant to Sections III or IV;

   and

   (b) those checks identify non-compliance with the rules on the protection of animals,

   the official auxiliary is immediately to inform the official veterinarian and, if necessary in cases of urgency, is to take the necessary measures referred to in paragraphs 1 to 4 pending the arrival of the official veterinarian.

CHAPTER V: DECISIONS CONCERNING MEAT

1. Meat is to be declared unfit for human consumption if it:

   (a) derives from animals that have not undergone ante-mortem inspection, except for hunted wild game;

   (b) derives from animals the offal of which has not undergone post-mortem inspection, unless otherwise provided for under this Regulation or Regulation (EC) No 853/2004;
(c) derives from animals which are dead before slaughter, stillborn, unborn or slaughtered under the age of seven days;

(d) results from the trimming of sticking points;

(e) derives from animals affected by animal diseases for which animal health rules are laid down in the Union legislation listed in Annex I to Council Directive 2002/99/EC (1), except if it is obtained in conformity with the specific requirements provided for in that legislation, unless otherwise provided for in Section IV;

(f) derives from animals affected by a generalised disease, such as generalised septicaemia, pyaemia, toxaemia or viraemia;

(g) is not in conformity with microbiological criteria laid down under Community legislation to determine whether food may be placed on the market;

(h) exhibits parasitic infestation, unless otherwise provided for in Section IV;

(i) contains residues or contaminants in excess of the levels laid down in Community legislation. Any overshooting of the relevant level should lead to additional analyses whenever appropriate;

(j) without prejudice to more specific Community legislation, derives from animals or carcases containing residues of forbidden substances or from animals that have been treated with forbidden substances;

(k) consists of the liver and kidneys of animals more than two years old from regions where implementation of plans approved in accordance with Article 5 of Directive 96/23/EC has revealed the generalised presence of heavy metals in the environment;

(l) has been treated illegally with decontaminating substances;

(m) has been treated illegally with ionising or UV-rays;

(n) contains foreign bodies (except, in the case of wild game, material used to hunt the animal);

(o) exceeds the maximum permitted radioactivity levels laid down under Community legislation;

(p) indicates patho-physiological changes, anomalies in consistency, insufficient bleeding (except for wild game) or organoleptic anomalies, in particular a pronounced sexual odour;

(q) derives from emaciated animals;

(r) contains specified risk material, except as provided for under Community legislation;

(s) shows soiling, faecal or other contamination;

(t) consists of blood that may constitute a risk to public or animal health owing to the health status of any animal from which it derives or contamination arising during the slaughter process;

(u) in the opinion of the official veterinarian, after examination of all the relevant information, it may constitute a risk to public or animal health or is for any other reason not suitable for human consumption.

2. The official veterinarian may impose requirements concerning the use of meat derived from animals:

(a) having undergone emergency slaughter outside the slaughterhouse; or

(b) if derived from flocks where a treatment of the meat will be applied in accordance with Part (E) of Annex II to Regulation (EC) No 2160/2003 before placing the meat on the market.

(1) OJ L 18, 23.1.2003, p. 11.
SECTION III: RESPONSIBILITIES AND FREQUENCY OF CONTROLS

CHAPTER I: OFFICIAL AUXILIARIES

Official auxiliaries may assist the official veterinarian with all tasks, subject to the following restrictions and to any specific rules laid down in Section IV:

1. in relation to auditing tasks, official auxiliaries may only collect information regarding good hygienic practices and HACCP-based procedures;

2. in relation to ante-mortem inspection and checks concerning the welfare of animals, official auxiliaries may only help with purely practical tasks which may include a preselection of animals with abnormalities;

3. in relation to post-mortem inspection, the official veterinarian must regularly check the work of official auxiliaries and, in the case of animals having undergone emergency slaughter outside the slaughterhouse, carry out the inspection personally.

CHAPTER II: FREQUENCY OF CONTROLS

1. The competent authority is to ensure that at least one official veterinarian is present:

   (a) in slaughterhouses, throughout both ante-mortem and post-mortem inspection;

   and

   (b) in game handling establishments, throughout post-mortem inspection.

2. However, the competent authority may adapt this approach in certain slaughterhouses and game handling establishments identified on the basis of a risk analysis and in accordance with criteria laid down in accordance with Article 18, point 3, if there are any. In such cases:

   (a) the official veterinarian need not be present at the time of ante-mortem inspection in the slaughterhouse if:

      (i) an official veterinarian or an approved veterinarian carried out ante-mortem inspection at the holding of provenance, checked the food chain information and communicated the results of the check to the official auxiliary at the slaughterhouse,

      (ii) the official auxiliary at the slaughterhouse is satisfied that the food chain information does not point to any possible problem for food safety and that the animal's general state of health and welfare is satisfactory,

      and

      (iii) the official veterinarian regularly satisfies himself/herself that the official auxiliary is carrying out such checks properly;

   (b) the official veterinarian need not be present at all times during post-mortem inspection if:

      (i) an official auxiliary carries out post-mortem inspection and puts aside meat with abnormalities and all other meat from the same animal,

      (ii) the official veterinarian subsequently inspects all such meat,

      and

      (iii) the official auxiliary documents his/her procedures and findings in a manner that allows the official veterinarian to be satisfied that standards are being met.
However, in the case of poultry and lagomorphs, the official auxiliary may discard meat with abnormalities and, subject to Section IV, the official veterinarian need not systematically inspect all such meat.

3. The flexibility provided for in paragraph 2 does not apply:

(a) to animals that have undergone emergency slaughter;

(b) to animals suspected of having a disease or condition that may adversely affect human health;

(c) to bovine animals from herds that have not been declared officially free of tuberculosis;

(d) to bovine, ovine and caprine animals from herds that have not been declared officially free of brucellosis;

(e) in the case of an outbreak of animal diseases for which animal health rules are laid down in Union legislation. This concerns animals susceptible to the particular disease in question that come from the particular region as defined in Article 2 of Council Directive 64/432/EEC (1);

(f) when stricter controls are necessary to take account of emerging diseases or particular OIE listed diseases.

4. In cutting plants, the competent authority is to ensure that an official veterinarian or an official auxiliary is present when meat is being worked on with a frequency appropriate to achieving the objectives of this Regulation.

CHAPTER III: INVOLVEMENT OF SLAUGHTERHOUSE STAFF

A. SPECIFIC TASKS CONCERNING THE PRODUCTION OF MEAT FROM POULTRY AND LAGOMORPHS

The Member States may permit slaughterhouse staff to take over the activities of the official auxiliaries in controlling the production of poultry and rabbit meat under the following conditions:

(a) Where the establishment has used good hygiene practice in accordance with Article 4(4) of this Regulation and the HACCP procedure for at least 12 months, the competent authority may authorise staff of the establishment to carry out tasks of official auxiliaries. This authorisation may only be granted if the staff of the establishment have been trained, to the satisfaction of the competent authority, in the same way as the official auxiliaries for the tasks of official auxiliaries or for the specific tasks they are authorised to perform. This staff must be placed under the supervision, direction and responsibility of the official veterinarian. In these circumstances, the official veterinarian shall be present at ante-mortem and post-mortem examinations, shall supervise these activities and carry out regular performance tests to ensure that the performance of the slaughterhouse staff meets the specific criteria laid down by the competent authority, and shall document the results of those performance tests. Where the level of hygiene of the establishment is affected by the work of this staff, where this staff does not carry out the tasks properly or where in general this staff carries out its work in a manner that the competent authority considers unsatisfactory, this staff shall be replaced by official auxiliaries.

(1) OJ 121, 29.7.1964, p. 1977/64.
(b) The competent authority of the Member State shall decide, in principle and on a case-by-case basis, whether to permit the implementation of the system described above. Where the Member State decides in principle in favour of this system, it shall inform the Commission of that decision and its associated conditions. For food business operators in a Member State implementing the system, the actual use of the system is optional. Food business operators shall not be forced by the competent authority to introduce the system described here. Where the competent authority is not convinced that the food business operator satisfies the requirements, the system shall not be implemented in that establishment. In order to assess this, the competent authority shall carry out an analysis of the production and inspection records, the type of activities undertaken in the establishment, the history of compliance with rules, the expertise, professional attitude and sense of responsibility of the slaughterhouse staff in regard to food safety, together with other relevant information.

B. SPECIFIC SAMPLING AND TESTING TASKS

Slaughterhouse staff who have received specific training, under the supervision of the official veterinarian, may, under the responsibility and the supervision of the official veterinarian, carry out specific sampling and testing tasks in respect of animals of all species.

CHAPTER IV: PROFESSIONAL QUALIFICATIONS

A. OFFICIAL VETERINARIANS

1. The competent authority may appoint only veterinarians who have passed a test meeting the requirements of paragraph 2 as official veterinarians.

2. The competent authority must make arrangements for the test. The test is to confirm knowledge of the following subjects to the extent necessary depending on the veterinarian's background and qualifications:

(a) national and Community legislation on veterinary public health, food safety, animal health, animal welfare and pharmaceutical substances;

(b) principles of the common agricultural policy, market measures, export refunds and fraud detection (including the global context: WTO, SPS, Codex Alimentarius, OIE);

(c) essentials of food processing and food technology;

(d) principles, concepts and methods of good manufacturing practice and quality management;

(e) pre-harvest quality management (good farming practices);

(f) promotion and use of food hygiene, food related safety (good hygiene practices);

(g) principles, concepts and methods of risk-analysis;

(h) principles, concepts and methods of HACCP, use of HACCP throughout the food production food chain;

(i) prevention and control of food-borne hazards related to human health;

(j) population dynamics of infection and intoxication;

(k) diagnostic epidemiology;

(l) monitoring and surveillance systems;
(m) auditing and regulatory assessment of food safety management systems;

(n) principles and diagnostic applications of modern testing methods;

(o) information and communication technology as related to veterinary public health;

(p) data-handling and applications of biostatistics;

(q) investigations of outbreaks of food-borne diseases in humans;

(r) relevant aspects concerning TSEs;

(s) animal welfare at the level of production, transport and slaughter;

(t) environmental issues related to food production (including waste management);

(u) precautionary principle and consumer concerns;

and

(v) principles of training of personnel working in the production chain.

Candidates may acquire the required knowledge as part of their basic veterinary training, or through training undertaken, or professional experience acquired, after qualifying as veterinarians. The competent authority may arrange for different tests to take account of candidates’ background. However, when the competent authority is satisfied that a candidate has acquired all the required knowledge as part of a university degree, or through continuing education resulting in a postgraduate qualification, it may waive the requirement for a test.

3. The veterinarian is to have aptitude for multidisciplinary cooperation.

4. In addition, each official veterinarian is to undergo practical training for a probationary period of at least 200 hours before starting to work independently. During this period the probationer is to work under the supervision of existing official veterinarians in slaughterhouses, cutting plants, inspection posts for fresh meat and on holdings. The training is to concern the auditing of food safety management systems in particular.

5. The official veterinarian is to maintain up-to-date knowledge and to keep abreast of new developments through regular continuing education activities and professional literature. The official veterinarian is, wherever possible, to undertake annual continuing education activities.

6. Veterinarians already appointed as official veterinarians must have adequate knowledge of the subjects mentioned in paragraph 2. Where necessary, they are to acquire this knowledge through continuing education activities. The competent authority is to make adequate provision in this regard.

7. Notwithstanding paragraphs 1 to 6, Member States may lay down specific rules for official veterinarians working on a part-time basis who are responsible for inspecting small businesses.
B. OFFICIAL AUXILIARIES

1. The competent authority may appoint as official auxiliaries only persons who have undergone training and passed a test in accordance with the following requirements.

2. The competent authority must make arrangements for such tests. To be eligible for these tests, candidates must prove that they have received:

(a) at least 500 hours of theoretical training and at least 400 hours of practical training, covering the areas specified in paragraph 5;

and

(b) such additional training as is required to enable official auxiliaries to undertake their duties competently.

3. The practical training referred to in paragraph 2(a) is to take place in slaughterhouses and cutting plants, under the supervision of an official veterinarian, and on holdings and in other relevant establishments.

4. Training and tests are to concern principally red meat or poultry meat. However, persons who undergo training for one of the two categories and passed the test need only undergo abridged training to pass the test for the other category. Training and test should cover wild game, farmed game and lagomorphs, where appropriate.

5. Training for official auxiliaries is to cover, and tests are to confirm knowledge of, the following subjects:

(a) in relation to holdings:

(i) theoretical part:

— familiarity with the farming industry organisation, production methods, international trade etc.,

— good livestock husbandry practices,

— basic knowledge of diseases, in particular zoonoses — viruses, bacteria, parasites etc.,

— monitoring for disease, use of medicines and vaccines, residue testing,

— hygiene and health inspection,

— animal welfare on the farm and during transport,

— environmental requirements — in buildings, on farms and in general,

— relevant laws, regulations and administrative provisions,

— consumer concerns and quality control;

(ii) practical part:

— visits to holdings of different types and using different rearing methods,

— visits to production establishments,

— observation of the loading and unloading of animals,
— laboratory demonstrations,
— veterinary checks,
— documentation;

(b) in relation to slaughterhouses and cutting plants:

(i) theoretical part:

— familiarity with the meat industry organisation, production methods, international trade and slaughter and cutting technology,
— basic knowledge of hygiene and good hygienic practices, and in particular industrial hygiene, slaughter, cutting and storage hygiene, hygiene of work,
— HACCP and the audit of HACCP-based procedures,
— animal welfare on unloading after transport and at the slaughterhouse,
— basic knowledge of the anatomy and physiology of slaughtered animals,
— basic knowledge of the pathology of slaughtered animals,
— basic knowledge of the pathological anatomy of slaughtered animals,
— relevant knowledge concerning TSEs and other important zoonoses and zoonotic agents,
— knowledge of methods and procedures for the slaughter, inspection, preparation, wrapping, packaging and transport of fresh meat,
— basic knowledge of microbiology,
— ante-mortem inspection,
— examination for trichinosis,
— post-mortem inspection,
— administrative tasks,
— knowledge of the relevant laws, regulations and administrative provisions,
— sampling procedure,
— fraud aspects;

(ii) practical part:

— animal identification,
— age checks,
— inspection and assessment of slaughtered animals,
— post-mortem inspection in a slaughterhouse,
— examination for trichinosis,
— identification of animal species by examination of typical parts of the animal,
— identifying and commenting on parts of slaughtered animals in which changes have occurred,
— hygiene control, including the audit of the good hygiene practices and the HACCP-based procedures,
— recording the results of ante-mortem inspection,
— sampling,
— traceability of meat,
— documentation.

6. Official auxiliaries are to maintain up-to-date knowledge and to keep abreast of new developments through regular continuing education activities and professional literature. The official auxiliary is, wherever possible, to undertake annual continuing education activities.

7. Persons already appointed as official auxiliaries must have adequate knowledge of the subjects mentioned in paragraph 5. Where necessary, they are to acquire this knowledge through continuing education activities. The competent authority is to make adequate provision in this regard.

8. However, when official auxiliaries carry out only sampling and analysis in connection with examinations for trichinosis, the competent authority need only ensure that they receive training appropriate to these tasks.

SECTION IV: SPECIFIC REQUIREMENTS

CHAPTER I: DOMESTIC BOVINE ANIMALS

A. BOVINE ANIMALS UNDER SIX WEEKS OLD

Carcasses and offal of bovine animals under six weeks old are to undergo the following post-mortem inspection procedures:

1. visual inspection of the head and throat; incision and examination of the retropharyngeal lymph nodes \((\text{Lnn retropharyngiales})\); inspection of the mouth and fauces; palpation of the tongue;

2. visual inspection of the lungs, trachea and oesophagus; palpation of the lungs; incision and examination of the bronchial and mediastinal lymph nodes \((\text{Lnn. bifurcations, eparteriales and mediastinales})\). The trachea and the main branches of the bronchi must be opened lengthwise and the lungs must be incised in their posterior third, perpendicular to their main axes; these incisions are not necessary where the lungs are excluded from human consumption;

3. visual inspection of the pericardium and heart, the latter being incised lengthwise so as to open the ventricles and cut through the interventricular septum;

4. visual inspection of the diaphragm;

5. visual inspection of the liver and the hepatic and pancreatic lymph nodes, \((\text{Lnn portales})\); palpation and, if necessary, incision of the liver and its lymph nodes;

6. visual inspection of the gastro-intestinal tract, the mesentery, the gastric and mesenteric lymph nodes \((\text{Lnn. gastrici, mesenterici, craniales and caudales})\); palpation and, if necessary, incision of the gastric and mesenteric lymph nodes;

7. visual inspection and, if necessary, palpation of the spleen;

8. visual inspection of the kidneys; incision, if necessary, of the kidneys and the renal lymph nodes \((\text{Lnn. renales})\);
9. visual inspection of the pleura and peritoneum;

10. visual inspection and palpation of the umbilical region and the joints. In the event of doubt, the umbilical region must be incised and the joints opened; the synovial fluid must be examined.

B. BOVINE ANIMALS OVER SIX WEEKS OLD

Carcasses and offal of bovine animals over six weeks old are to undergo the following post-mortem inspection procedures:

1. visual inspection of the head and throat; incision and examination of the sub-maxillary, retropharyngeal and parotid lymph nodes (Lnn retropharyngiales, mandibulares and parotidei); examination of the external masseters, in which two incisions must be made parallel to the mandible, and the internal masseters (internal pterygoid muscles), which must be incised along one plane. The tongue must be freed to permit a detailed visual inspection of the mouth and the fauces and must itself be visually inspected and palpated; ►M4◄

2. inspection of the trachea and oesophagus; visual examination and palpation of the lungs; incision and examination of the bronchial and mediastinal lymph nodes (Lnn. bifurcations, eparteriales and mediastinales). The trachea and the main branches of the bronchi must be opened lengthways and the lungs must be incised in their posterior third, perpendicular to their main axes; these incisions are not necessary where the lungs are excluded from human consumption;

3. visual inspection of the pericardium and heart, the latter being incised lengthways so as to open the ventricles and cut through the interventricular septum;

4. visual inspection of the diaphragm;

5. visual inspection and palpation of the liver and the hepatic and pancreatic lymph nodes, (Lnn portales); incision of the gastric surface of the liver and at the base of the caudate lobe to examine the bile ducts;

6. visual inspection of the gastro-intestinal tract, the mesentery, the gastric and mesenteric lymph nodes (Lnn. gastrici, mesenterici, craniales and caudales); palpation and, if necessary, incision of the gastric and mesenteric lymph nodes;

7. visual inspection and, if necessary, palpation of the spleen;

8. visual inspection of the kidneys and incision, if necessary, of the kidneys and the renal lymph nodes (Lnn. renales);

9. visual inspection of the pleura and the peritoneum;

10. visual inspection of the genital organs (except for the penis, if already discarded);

11. visual inspection and, if necessary, palpation and incision of the udder and its lymph nodes (Lnn. supramammarii). In cows, each half of the udder must be opened by a long, deep incision as far as the lactiferous sinuses (sinus lactiferes) and the lymph nodes of the udder must be incised, except when the udder is excluded from human consumption.
CHAPTER II: DOMESTIC SHEEP AND GOATS

Carcasses and offal of sheep and goats are to undergo the following post-mortem inspection procedures:

1. visual inspection of the head after flaying and, in the event of doubt, examination of the throat, mouth, tongue and retropharyngeal and parotid lymph nodes. Without prejudice to animal-health rules, these examinations are not necessary if the competent authority is able to guarantee that the head, including the tongue and the brains, will be excluded from human consumption;

2. visual inspection of the lungs, trachea and oesophagus; palpation of the lungs and the bronchial and mediastinal lymph nodes (Lnn. bifucationes, eparteriales and mediastinales); in the event of doubt, these organs and lymph nodes must be incised and examined;

3. visual inspection of the pericardium and heart; in the event of doubt, the heart must be incised and examined;

4. visual inspection of the diaphragm;

5. visual inspection of the liver and the hepatic and pancreatic lymph nodes, (Lnn. portales); palpation of the liver and its lymph nodes; incision of the gastric surface of the liver to examine the bile ducts;

6. visual inspection of the gastro-intestinal tract, the mesentery and the gastric and mesenteric lymph nodes (Lnn. gastrici, mesenterici, craniales and caudales);

7. visual inspection and, if necessary, palpation of the spleen;

8. visual inspection of the kidneys; incision, if necessary, of the kidneys and the renal lymph nodes (Lnn. renales);

9. visual inspection of the pleura and peritoneum;

10. visual inspection of the genital organs (except for the penis, if already discarded);

11. visual inspection of the udder and its lymph nodes;

12. visual inspection and palpation of the umbilical region and joints of young animals. In the event of doubt, the umbilical region must be incised and the joints opened; the synovial fluid must be examined.

CHAPTER III: DOMESTIC SOLIPEDS

Carcasses and offal of solipeds are to undergo the following post-mortem inspection procedures:

1. visual inspection of the head and, after freeing the tongue, the throat; palpation and, if necessary, incision of the sub-maxillary, retropharyngeal and parotid lymph nodes (Lnn retropharyngiales, mandibulares and parotidei). The tongue must be freed to permit a detailed visual inspection of the mouth and the fauces and must itself be visually examined and palpated;

2. visual inspection of the lungs, trachea and oesophagus; palpation of the lungs; palpation and, if necessary, incision of the bronchial and mediastinal lymph nodes (Lnn. bifucationes, eparteriales and mediastinales). The trachea and the main branches of the bronchi must be opened lengthwise and the lungs must be incised in their posterior third, perpendicular to their main axes; however, these incisions are not necessary where the lungs are excluded from human consumption;
3. visual inspection of the pericardium and the heart, the latter being incised lengthwise so as to open the ventricles and cut through the interventricular septum;

4. visual inspection of the diaphragm;

5. visual inspection, palpation and, if necessary, incision of the liver and the hepatic and pancreatic lymph nodes, (Lnn. portales);

6. visual inspection of the gastro-intestinal tract, the mesentery and the gastric and mesenteric lymph nodes (Lnn. gastrici, mesenterici, craniales and caudales); incision, if necessary, of the gastric and mesenteric lymph nodes;

7. visual inspection and, if necessary, palpation of the spleen;

8. visual inspection and palpation of the kidneys; incision, if necessary, of the kidneys and the renal lymph nodes (Lnn. renales);

9. visual inspection of the pleura and peritoneum;

10. visual inspection of the genital organs of stallions (except for the penis, if already discarded) and mares;

11. visual inspection of the udder and its lymph nodes (Lnn. supramammarii) and, if necessary, incision of the supramammary lymph nodes;

12. visual inspection and palpation of the umbilical region and joints of young animals. In the event of doubt, the umbilical region must be incised and the joints opened; the synovial fluid must be examined;

13. all grey or white horses must be inspected for melanosis and melanomata by examination of the muscles and lymph nodes (Lnn. subrhomboidei) of the shoulders beneath the scapular cartilage after loosening the attachment of one shoulder. The kidneys must be exposed and examined by incision through the entire kidney.

CHAPTER IV: DOMESTIC SWINE
A. ANTE-MORTEM INSPECTION

1. The competent authority may decide that pigs intended for slaughter are to be submitted to ante-mortem inspection at the holding of provenance. In that case, slaughter of a lot of pigs from a holding may be authorised only if:

   (a) the health certificate provided for in Chapter X, Part A, accompanies them;

   and

   (b) the requirements of paragraphs 2 to 5 are complied with.

2. Ante-mortem inspection at the holding of provenance is to comprise:

   (a) checks on records or documentation at the holding, including food chain information;

   (b) the examination of the pigs to determine whether:

      (i) they have a disease or condition which may be transmitted to animals or humans through handling or eating the meat, or are behaving, individually or collectively, in a manner indicating that such a disease may occur,
(ii) they show disturbance of general behaviour or signs of disease which may make the meat unfit for human consumption,

or

(iii) there is evidence or reasons to suspect that they may contain chemical residues in excess of the levels laid down in Community legislation, or residues of forbidden substances.

3. An official veterinarian or an approved veterinarian is to carry out ante-mortem inspection at the holding. The pigs are to be sent directly to slaughter and not to be mixed with other pigs.

4. Ante-mortem inspection at the slaughterhouse need cover only:

(a) a control of the animals’ identification;

and

(b) a screening to ascertain whether animal welfare rules have been complied with and whether signs of any condition which might adversely affect human or animal health are present. An official auxiliary may carry out this screening.

5. When pigs are not slaughtered within three days of the issue of the health certificate provided for in paragraph 1(a):

(a) if the pigs have not left the holding of provenance for the slaughterhouse, they are to be re-examined and a new health certificate issued;

(b) if the pigs are already en route for or at the slaughterhouse, slaughter may be authorised once the reason for the delay has been assessed, provided that the pigs undergo a further veterinary ante-mortem inspection.

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B. POST-MORTEM INSPECTION

1. Carcases and offal of pigs are to undergo the following post-mortem inspection procedures:

(a) visual inspection of the head and throat; visual inspection of the mouth, fauces and tongue;

(b) visual inspection of the lungs, trachea and oesophagus;

(c) visual inspection of the pericardium and heart;

(d) visual inspection of the diaphragm;

(e) visual inspection of the liver and the hepatic and pancreatic lymph nodes (Lnn. portales);
(f) visual inspection of the gastro-intestinal tract, the mesentery, the
gastric and mesenteric lymph nodes (Lnn. gastrici, mesenterici,
craniales and caudales);

(g) visual inspection of the spleen;

(h) visual inspection of the kidneys;

(i) visual inspection of the pleura and peritoneum;

(j) visual inspection of the genital organs (except for the penis, if already
discarded);

(k) visual inspection of the udder and its lymph nodes (Lnn. supram-
annarii);

(l) visual inspection of the umbilical region and joints of young animals.

2. The official veterinarian shall proceed with additional post-mortem
inspection procedures using incision and palpation of the carcase and
offal, where, in his or her opinion, one of the following indicates a
possible risk to public health, animal health or animal welfare:

(a) the checks and analysis of the food chain information carried out in
accordance with Part A of Chapter II of Section I;

(b) the findings of the ante-mortem inspection carried out in accordance
with Part B of Chapter II of Section I and Part A of this Chapter;

(c) the results of the verifications concerning compliance with animal
welfare rules carried out in accordance with Part C of Chapter II of
Section I;

(d) the findings of post-mortem inspection carried out in accordance with
Part D of Chapter II of Section I and point 1 of this part;

(e) additional epidemiological data or other data from the holding of
provenance of the animals.

3. Depending on the identified risks, the additional post-mortem procedures
referred to in point 2 may include:

(a) incision and examination of the submaxillary lymph nodes (Lnn.
mandibulares);

(b) palpation of the lungs and the bronchial and mediastinal lymph nodes
(Lnn. bifurciones, eparteriales and mediastinales). The trachea and
the main branches of the bronchi must be opened lengthwise and the
lungs must be incised in their posterior third, perpendicular to their
main axes; those incisions are not necessary where the lungs are
excluded from human consumption;

(c) incision of the heart lengthwise so as to open the ventricles and cut
through the interventricular septum;

(d) palpation of the liver and its lymph nodes;

(e) palpation and, if necessary, incision of the gastric and mesenteric
lymph nodes;
(f) palpation of the spleen;

(g) incision of the kidneys and the renal lymph nodes (Lnn. renales);

(h) incision of the supramammary lymph nodes;

(i) palpation of the umbilical region and joints of young animals and, if necessary, incision of the umbilical region and opening of the joints.

CHAPTER V: POULTRY

A. ANTE-MORTEM INSPECTION

1. The competent authority may decide that poultry intended for slaughter are to be submitted to ante-mortem inspection at the holding of provenance. In that case, slaughter of a flock of birds from a holding may be authorised only if:

(a) the health certificate provided for in Chapter X, Part A, accompanies them;

and

(b) the requirements of paragraphs 2 to 5 are complied with.

2. Ante-mortem inspection on the holding of provenance is to comprise:

(a) checks on records or documentation at the holding, including food chain information;

(b) a flock inspection, to determine whether the birds:

   (i) have a disease or condition which may be transmitted to animals or humans through handling or eating the meat, or are behaving in a manner indicating that such a disease may occur,

   (ii) show disturbance of general behaviour or signs of disease which may make the meat unfit for human consumption,

   or

   (iii) show evidence that they may contain chemical residues in excess of the levels laid down in Community legislation, or residues of forbidden substances.

3. An official veterinarian or an approved veterinarian is to carry out ante-mortem inspection at the holding.
4. Ante-mortem inspection at the slaughterhouse need only cover:

(a) a control of the animals’ identification;

and

(b) a screening to ascertain whether animal welfare rules have been complied with and whether signs of any condition which might adversely affect human or animal health are present. An official auxiliary may carry out this screening.

5. When birds are not slaughtered within three days of the issue of the health certificate referred to in paragraph I(a):

(a) if the flock has not left the holding of provenance for the slaughterhouse, it is to be re-examined and a new health certificate issued;

(b) if the flock is already en route for or at the slaughterhouse, slaughter may be authorised once the reason for the delay has been assessed, provided that the flock is re-examined.

6. When ante-mortem inspection is not carried out at the holding, the official veterinarian is to carry out a flock inspection at the slaughterhouse.

7. If the birds show clinical symptoms of a disease, they may not be slaughtered for human consumption. However, killing of these birds on the slaughter line may take place at the end of the normal slaughter process, if precautions are taken to avoid the risk of spreading pathogenic organisms and to clean and disinfect the facilities immediately after killing.

8. In the case of poultry reared for the production of ‘foie gras’ and delayed eviscerated poultry slaughtered at the holding of provenance, ante-mortem inspection is to be carried out in accordance with paragraphs 2 and 3. A certificate conforming to the model set out in Part C is to accompany the uneviscerated carcases to the slaughterhouse or cutting plant.

B. Post-mortem inspection

1. All birds are to undergo post-mortem inspection in accordance with Sections I and III. In addition, the official veterinarian is personally to carry out the following checks:

(a) daily inspection of the viscera and body cavities of a representative sample of birds;

(b) a detailed inspection of a random sample, from each batch of birds having the same origin, of parts of birds or entire birds declared unfit for human consumption following post-mortem inspection;

and

(c) any further investigations necessary when there is reason to suspect that the meat from the birds concerned could be unfit for human consumption.

2. In the case of poultry reared for the production of ‘foie gras’ and delayed eviscerated poultry obtained at the holding of provenance, post-mortem inspection is to include a check on the certificate accompanying the carcases. When such carcases are transported directly from the holding to a cutting plant, post-mortem inspection is to take place at the cutting plant.
C. SPECIMEN HEALTH CERTIFICATE

HEALTH CERTIFICATE

for poultry intended for the production of foie gras and delayed eviscerated
poultry slaughtered at the holding of provenance

Competent service: ........................................................................................................

No: .................................................................................................................................

1. Identification of uneviscerated carcases

Species: ............................................................................................................................

Number: .........................................................................................................................

2. Provenance of uneviscerated carcases

Address of holding: ........................................................................................................

3. Destination of uneviscerated carcases

The uneviscerated carcases will be transported to the following cutting plant: ..............

.................................................................................................................................

4. Declaration

I, the undersigned, declare that:

— the uneviscerated carcases described above are of birds which were examined before slaughter on the
  abovementioned holding at ........... (time) on .......... (date) and found to be healthy;

— the records and documentation concerning these animals satisfied the legal requirements and do not prohibit
  slaughter of the birds.

Done at: ..........................................................................................................................

(Place)

on: ...............................................................................................................................

(Date)

Stamp

.................................................................................................................................

(Signature of the official or approved veterinarian)
CHAPTER VI: FARMED LAGOMORPHS
The requirements for poultry are to apply to farmed lagomorphs.

CHAPTER VII: FARMED GAME

A. Ante-mortem inspection
1. Ante-mortem inspection may be carried out at the holding of provenance when the requirements of Annex III, Section III, to Regulation (EC) No 853/2004 are satisfied. In this case, an official veterinarian or an approved veterinarian is to carry out ante-mortem inspection.
2. Ante-mortem inspection at the holding is to include checks on the records or documentation at the holding, including food chain information.
3. When ante-mortem inspection takes place no more than three days before the arrival of the animals at the slaughterhouse, and animals are delivered to the slaughterhouse live, ante-mortem inspection at the slaughterhouse need only cover:
   (a) a control of the animals' identification;
   and
   (b) a screening to ascertain whether animal welfare rules have been complied with and whether signs of any condition which might adversely affect human or animal health are present.
4. A certificate conforming to the specimen in Chapter X, Part A, is to accompany live animals inspected at the holding. A certificate conforming to the specimen in Chapter X, Part B, is to accompany animals inspected and slaughtered at the holding. A certificate conforming to the specimen in Chapter X, Part C, is to accompany animals inspected and slaughtered at the holding in accordance with point 3a of Section III of Annex III to Regulation (EC) No 853/2004.
5. When the competent authority authorises that the food business operator may attest the correct slaughter and bleeding of animals, the official veterinarian or approved veterinarian shall carry out regular checks on the performance of the person carrying out the slaughter and bleeding.

B. Post-mortem inspection
1. This inspection is to include palpation and, where judged necessary, incision of those parts of the animal which have undergone any change or are suspect for any other reason.
2. Post-mortem inspection procedures described for bovine and ovine animals, domestic swine and poultry are to be applied to the corresponding species of farmed game.
3. When the animals have been slaughtered at the holding, the official veterinarian at the slaughterhouse is to check the certificate accompanying them.

CHAPTER VIII: WILD GAME

A. Post-mortem inspection
1. Wild game is to be inspected as soon as possible after admission to the game handling establishment.
2. The official veterinarian is to take account of the declaration or information that the trained person involved in hunting the animal has provided in accordance with Regulation (EC) No 853/2004.

2a. The official veterinarian is to check that a health certificate conforming to the specimen set out in the Annex to Commission Implementing Regulation (EU) No 636/2014 (1) or the declaration(s) accompanies the unskinned large wild game transported to the game-handling establishment from the territory of another Member State, in accordance with point 8(b) of Chapter II of Section IV of Annex III to Regulation (EC) No 853/2004. The official veterinarian is to take into account the content of that certificate or declaration(s).

3. During post-mortem inspection, the official veterinarian is to carry out:

(a) a visual examination of the carcase, its cavities and, where appropriate, organs with a view to:

(i) detecting any abnormalities not resulting from the hunting process. For this purpose, the diagnosis may be based on any information that the trained person has provided concerning the behaviour of the animal before killing,

(ii) checking that death was not caused by reasons other than hunting.

If an assessment cannot be made on the basis of visual examination alone, a more extensive inspection must be carried out in a laboratory;

(b) an investigation of organoleptic abnormalities;

(c) palpation of organs, where appropriate;

(d) where there are serious grounds for suspecting the presence of residues or contaminants, an analysis by sampling of residues not resulting from the hunting process, including environmental contaminants. When a more extensive inspection is made on the basis of such suspicions, the veterinarian must wait until that inspection has been concluded before assessing all the game killed during a specific hunt, or those parts suspected of showing the same abnormalities;

(e) examination for characteristics indicating that the meat presents a health risk, including:

(i) abnormal behaviour or disturbance of the general condition of the live animal, as reported by the hunter,

(ii) the generalised presence of tumours or abscesses affecting different internal organs or muscles,

(iii) arthritis, orchitis, pathological changes in the liver or the spleen, inflammation of the intestines or the umbilical region,

(iv) the presence of foreign bodies not resulting from the hunting process in the body cavities, stomach or intestines or in the urine, where the pleura or peritoneum are discoloured (when relevant viscera are present),

(v) the presence of parasites,

(vi) formation of a significant amount of gas in the gastro-intestinal tract with discolouring of the internal organs (when these viscera are present),

(vii) significant abnormalities of colour, consistency or odour of muscle tissue or organs,

(viii) aged open fractures,

(ix) emaciation and/or general or localised oedema,

(x) recent pleural or peritoneal adhesions,

and

(xi) other obvious extensive changes, such as putrefaction.

4. Where the official veterinarian so requires, the vertebral column and the head are to be split lengthwise.
5. In the case of small wild game not eviscerated immediately after killing, the official veterinarian is to carry out a post-mortem inspection on a representative sample of animals from the same source. Where inspection reveals a disease transmissible to man or any of the characteristics listed in paragraph 3(e), the official veterinarian is to carry out more checks on the entire batch to determine whether it must be declared unfit for human consumption or whether each carcass must be inspected individually.

6. In the event of doubt, the official veterinarian may perform any further cuts and inspections of the relevant parts of the animals necessary to reach a final diagnosis.

B. Decisions following controls

In addition to the cases provided for in Section II, Chapter V, meat presenting during post-mortem inspection any of the characteristics listed in paragraph 3(e) of Part A is to be declared unfit for human consumption.

CHAPTER IX: SPECIFIC HAZARDS

A. Transmissible spongiform encephalopathies

Official controls carried out in relation to TSEs are to take account of the requirements of Regulation (EC) No 999/2001 and other relevant Community legislation.

B. Cysticercosis

1. The post-mortem inspection procedures described in Chapters I and IV are the minimum requirements for the examination for cysticercosis in bovine animals over six weeks old and swine. In addition, specific serological tests may be used. In the case of bovines over six weeks old, incision of the masseters at post-mortem inspection is not compulsory when a specific serological test is used. The same applies when bovine animals over six weeks old have been raised on a holding officially certified to be free of cysticercosis.

2. Meat infected with cysticercus is to be declared unfit for human consumption. However, when the animal is not generally infected with cysticercus, the parts not infected may be declared fit for human consumption after having undergone a cold treatment.

C. Trichinosis

1. Carcases of swine (domestic, farmed game and wild game), solipeds and other species susceptible to trichinosis are to be examined for trichinosis in accordance with applicable Community legislation, unless that legislation provides otherwise.

2. Meat from animals infected with trichinae is to be declared unfit for human consumption.

D. Glanders

1. Where appropriate, solipeds are to be examined for glanders. Examination for glanders in solipeds is to include a careful examination of mucous membranes from the trachea, larynx, nasal cavities and sinuses and their ramifications, after splitting the head in the median plane and excising the nasal septum.

2. Meat from horses in which glanders has been diagnosed are to be declared unfit for human consumption.
E. Tuberculosis

1. When animals have reacted positively or inconclusively to tuberculin, or there are other grounds for suspecting infection, they are to be slaughtered separately from other animals, taking precautions to avoid the risk of contamination of other carcases, the slaughter line and staff present in the slaughterhouse.

2. All meat from animals in which post-mortem inspection has revealed localised tuberculous lesions in a number of organs or a number of areas of the carcase is to be declared unfit for human consumption. However, when a tuberculous lesion has been found in the lymph nodes of only one organ or part of the carcase, only the affected organ or part of the carcase and the associated lymph nodes need be declared unfit for human consumption.

F. Brucellosis

1. When animals have reacted positively or inconclusively to a brucellosis test, or there are other grounds for suspecting infection, they are to be slaughtered separately from other animals, taking precautions to avoid the risk of contamination of other carcases, the slaughter line and staff present in the slaughterhouse.

2. Meat from animals in which post-mortem inspection has revealed lesions indicating acute infection with brucellosis is to be declared unfit for human consumption. In the case of animals reacting positively or inconclusively to a brucellosis test, the udder, genital tract and blood must be declared unfit for human consumption even if no such lesion is found.

G. Salmonella

1. Without prejudice to the first paragraph of Article 1 to Commission Regulation (EC) No 2073/2005 (1), the competent authority shall verify the correct implementation by food business operators of the point 2.1.4 (process hygiene criterion for Salmonella on pig carcases) of Annex I to that Regulation by applying the following measures:

   (a) official sampling using the same method and sampling area as food business operators. At least 49 (2) random samples shall be taken in each slaughterhouse each year. This number of samples may be reduced in small slaughterhouses based on a risk evaluation; and/or

   (b) collecting all information on the total number and the number of Salmonella positive samples taken by food business operators in accordance with Article 5(5) of Regulation (EC) No 2073/2005, within the frame of point 2.1.4 of Annex I thereof; and/or

   (c) collecting all information on the total number and the number of Salmonella positive samples taken within the frame of national control programmes in Member States or regions of Member States for which special guarantees have been approved in accordance with Article 8 of Regulation (EC) No 853/2004 as regards pork production;

2. If the process hygiene criterion is not complied with at several occasions, the competent authority shall require an action plan from the food business operator concerned and strictly supervise its outcome.

3. The total number and the number of Salmonella positive samples, differentiating between samples taken under (a), (b) and (c), when applied, shall be reported in accordance with Article 9(1) of Directive 2003/99/EC of the European Parliament and of the Council (3).

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(2) If all negative, 95 % statistical certainty is provided that the prevalence is below 6 %.
CHAPTER X: SPECIMEN HEALTH CERTIFICATE

A. SPECIMEN HEALTH CERTIFICATE FOR LIVE ANIMALS

HEALTH CERTIFICATE

for live animals transported from the holding to the slaughterhouse

Competent service: ..............................................................................................................................

No: .................................................................................................................................................

1. Identification of the animals
   Species: .........................................................................................................................................
   Number of animals: ........................................................................................................................
   Identification marking: ....................................................................................................................

2. Provenance of the animals
   Address of holding of provenance: ............................................................................................
   Identification of house (?): ..........................................................................................................

3. Destination of the animals
   The animals will be transported to the following slaughterhouse: ..............................................
   by the following means of transport: ...........................................................................................

4. Other relevant information
   ......................................................................................................................................................

5. Declaration
   I, the undersigned, declare that:
   — the animals described above were examined before slaughter at the abovementioned holding at .......... (time) on .......... (date) and were found to be healthy,
   — the records and documentation concerning these animals satisfied the legal requirements and do not prohibit
     slaughter of the animals.
   Done at: ........................................................................................................................................
   (Place)
   on: ..............................................................................................................................................
   (Date)

Stamp

........................................................................................................................................................
(Signature of official or approved veterinarian)

(*) optional
HEALTH CERTIFICATE

for animals slaughtered at the holding

Competent service: .................................................................................................................................

No: ..............................................................................................................................................................

1. Identification of the animals
Species: .............................................................................................................................................................
Number of animals: ...........................................................................................................................................
Identification marking: .........................................................................................................................................

2. Provenance of the animals
Address of holding of provenance: ........................................................................................................................
Identification of house (*): ...................................................................................................................................

3. Destination of the animals
The animals will be transported to the following slaughterhouse: ...........................................................................
..............................................................................................................................................................................
by the following means of transport: ...........................................................................................................................

4. Other relevant information
...........................................................................................................................................................................

5. Declaration
I, the undersigned, declare that:
— the animals described above were examined before slaughter at the abovementioned holding at .......... (time) on ........... (date) and were found to be healthy,
— they were slaughtered at the holding at ........... (time) on ........... (date) and slaughter and bleeding were carried out correctly,
— the records and documentation concerning these animals satisfied the legal requirements and did not prohibit slaughter of the animals.

Done at: ................................................................. (Place)
on: ................................................................. (Date)
Stamp

(Signature of official or approved veterinarian)

(*) optional
C. SPECIMEN HEALTH CERTIFICATE FOR FARMED GAME
SLAUGHTERED AT THE HOLDING in accordance with point 3a of

HEALTH CERTIFICATE

for farmed game slaughtered at the holding in accordance with point 3a of Section III of Annex III to
Regulation (EC) No 853/2004

Competent service: .................................................................................................................................................

No: ...........................................................................................................................................................................

1. Identification of the animals

Species: ........................................................................................................................................................................

Number of animals: ..................................................................................................................................................

Identification marking: ...............................................................................................................................................

2. Provenance of the animals

Address of holding of provenance: ..............................................................................................................................

Identification of house (*): ...........................................................................................................................................

3. Destination of the animals

The animals will be transported to the following slaughterhouse: .................................................................

...........................................................................................................................................................................

by the following means of transport: ........................................................................................................................

4. Other relevant information

...........................................................................................................................................................................

5. Declaration

I, the undersigned, declare that:

— the animals described above were examined before slaughter at the abovementioned holding at .................
  (time) on ................. (date) and were found to be healthy,

— the records and documentation concerning these animals satisfied the legal requirements and did not prohibit
  slaughter of the animals.

Done at: ...................................................................................................................................................................

(Place)

on: ..............................................................................................................................................................................

(Date)

Stamp

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(Signature of official or approved veterinarian)

(*) optional.
CHAPTER I: SCOPE

This Annex applies to live bivalve molluscs and, by analogy, to live echinoderms, live tunicates and live marine gastropods.

CHAPTER II: OFFICIAL CONTROLS CONCERNING LIVE BIVALVE MOLLUSCS FROM CLASSIFIED PRODUCTION AREAS

The reference method for analysis of E. coli is the detection and Most Probable Number (MPN) technique specified in EN/ISO 16649-3. Alternative methods may be used if they are validated against this reference method in accordance with the criteria in EN/ISO 16140.

A. CLASSIFICATION OF PRODUCTION AND RELAYING AREAS

1. The competent authority must fix the location and boundaries of production and relaying areas that it classifies. It may, where appropriate, do so in cooperation with the food business operator.

2. The competent authority must classify production areas from which it authorises the harvesting of live bivalve molluscs as being of one of three categories according to the level of faecal contamination. It may, where appropriate, do so in cooperation with the food business operator. In order to classify production areas, the competent authority must define a review period for sampling data from each production and relaying area in order to determine compliance with the standards referred to in this paragraph and in paragraphs 3, 4 and 5.

3. The competent authority may classify as being of Class A areas from which live bivalve molluscs may be collected for direct human consumption. Live bivalve molluscs placed on the market from these areas must meet the health standards laid down in Annex III, Section VII, Chapter V, of Regulation (EC) No 853/2004.

Samples of live bivalve molluscs from these areas must not exceed, in 80 % of samples collected during the review period, 230 E. coli per 100 g of flesh and intravalvular liquid. The remaining 20 % of samples must not exceed 700 E. coli per 100 g of flesh and intravalvular liquid.

When evaluating the results for the defined review period for maintenance of a Class A area, the competent authority can, based on a risk assessment on the basis of an investigation, decide to disregard an anomalous result exceeding the level of 700 E. coli per 100 g of flesh and intravalvular liquid.

4. The competent authority may classify as being of Class B areas from which live bivalve molluscs may be collected and only placed on the market for human consumption after treatment in a purification centre or after relaying so as to meet the health standards referred to in paragraph 3. Live bivalve molluscs from these areas must not exceed, in 90 % of the samples, 4 600 E. coli per 100 g of flesh and intravalvular liquid. In the remaining 10 % of samples, live bivalve molluscs must not exceed 46 000 E. coli per 100 g of flesh and intravalvular liquid.
5. The competent authority may classify as being of Class C areas from which live bivalve molluscs may be collected and only placed on the market after relaying over a long period so as to meet the health standards referred to in paragraph 3. Live bivalve molluscs from these areas must not exceed 46 000 E. coli per 100 g of flesh and intravalvular liquid.

6. If the competent authority decides in principle to classify a production or relaying area, it must:

(a) make an inventory of the sources of pollution of human or animal origin likely to be a source of contamination for the production area;

(b) examine the quantities of organic pollutants which are released during the different periods of the year, according to the seasonal variations of both human and animal populations in the catchment area, rainfall readings, waste-water treatment, etc.;

(c) determine the characteristics of the circulation of pollutants by virtue of current patterns, bathymetry and the tidal cycle in the production area;

and

(d) establish a sampling programme of bivalve molluscs in the production area which is based on the examination of established data, and with a number of samples, a geographical distribution of the sampling points and a sampling frequency which must ensure that the results of the analysis are as representative as possible for the area considered.

B. MONITORING OF CLASSIFIED RELAYING AND PRODUCTION AREAS

1. Classified relaying and production areas must be periodically monitored to check:

(a) that there is no malpractice with regard to the origin, provenance and destination of live bivalve molluscs;

(b) the microbiological quality of live bivalve molluscs in relation to the production and relaying areas;

(c) for the presence of toxin-producing plankton in production and relaying waters and biotoxins in live bivalve molluscs;

and

(d) for the presence of chemical contaminants in live bivalve molluscs.

2. To implement paragraph 1(b), (c) and (d), sampling plans must be drawn up providing for such checks to take place at regular intervals, or on a case-by-case basis if harvesting periods are irregular. The geographical distribution of the sampling points and the sampling frequency must ensure that the results of the analysis are as representative as possible for the area considered.

3. Sampling plans to check the microbiological quality of live bivalve molluscs must take particular account of:

(a) the likely variation in faecal contamination,

and

(b) the parameters referred to in paragraph 6 of Part A.

4. Sampling plans to check for the presence of toxin-producing plankton in production and relaying waters and for biotoxins in live bivalve molluscs must take particular account of possible variations in the presence of plankton containing marine biotoxins. Sampling must comprise:
(a) periodic sampling to detect changes in the composition of plankton containing toxins and their geographical distribution. Results suggesting an accumulation of toxins in mollusc flesh must be followed by intensive sampling;

(b) periodic toxicity tests using those molluscs from the affected area most susceptible to contamination.

5. The sampling frequency for toxin analysis in the molluscs is, as a general rule, to be weekly during the periods at which harvesting is allowed. This frequency may be reduced in specific areas, or for specific types of molluscs, if a risk assessment on toxins or phytoplankton occurrence suggests a very low risk of toxic episodes. It is to be increased where such an assessment suggests that weekly sampling would not be sufficient. The risk assessment is to be periodically reviewed in order to assess the risk of toxins occurring in the live bivalve molluscs from these areas.

6. When knowledge of toxin accumulation rates is available for a group of species growing in the same area, a species with the highest rate may be used as an indicator species. This will allow the exploitation of all species in the group if toxin levels in the indicator species are below the regulatory limits. When toxin levels in the indicator species are above the regulatory limits, harvesting of the other species is only to be allowed if further analysis on the other species shows toxin levels below the limits.

7. With regard to the monitoring of plankton, the samples are to be representative of the water column and to provide information on the presence of toxic species as well as on population trends. If any changes in toxic populations that may lead to toxin accumulation are detected, the sampling frequency of molluscs is to be increased or precautionary closures of the areas are to be established until results of toxin analysis are obtained.

8. Sampling plans to check for the presence of chemical contaminants must enable the detection of any overshooting of the levels laid down in Commission Regulation (EC) No 466/2001 (1).

C. DECISIONS AFTER MONITORING

1. Where the results of sampling show that the health standards for molluscs are exceeded, or that there may be otherwise a risk to human health, the competent authority must close the production area concerned, preventing the harvesting of live bivalve molluscs. However, the competent authority may reclassify a production area as being of Class B or C if it meets the relevant criteria set out in Part A and presents no other risk to human health.

2. The competent authority may re-open a closed production area only if the health standards for molluscs once again comply with Community legislation. If the competent authority closes a production because of the presence of plankton or excessive levels of toxins in molluscs, at least two consecutive results below the regulatory limit separated at least 48 hours are necessary to re-open it. The competent authority may take account of information on phytoplankton trends when taking this decision. When there are robust data on the dynamic of the toxicity for a given area, and provided that recent data on decreasing trends of toxicity are available, the competent authority may decide to re-open the area with results below the regulatory limit obtained from one single sampling.

D. ADDITIONAL MONITORING REQUIREMENTS

1. The competent authority is to monitor classified production areas from which it has forbidden the harvesting of bivalve molluscs or subjected harvesting to special conditions, to ensure that products harmful to human health are not placed on the market.

2. In addition to the monitoring of relaying and production zones referred to in paragraph 1 of Part B, a control system must be set up comprising laboratory tests to verify food business operators' compliance with the requirements for the end product at all stages of production, processing and distribution. This control system is, in particular, to verify that the levels of marine biotoxins and contaminants do not exceed safety limits and that the microbiological quality of the molluscs does not constitute a hazard to human health.

E. RECORDING AND EXCHANGE OF INFORMATION

The competent authority must:

(a) establish and keep up to date a list of approved production and relaying areas, with details of their location and boundaries, as well as the class in which the area is classified, from which live bivalve molluscs may be taken in accordance with the requirements of this Annex. This list must be communicated to interested parties affected by this Annex, such as producers, gatherers and operators of purification centres and dispatch centres;

(b) immediately inform the interested parties affected by this Annex, such as producers, gatherers and operators of purification centres and dispatch centres, about any change of the location, boundaries or class of a production area, or its closure, be it temporary or final;

and

(c) act promptly where the controls prescribed in this Annex indicate that a production area must be closed or reclassified or can be re-opened.

F. FOOD BUSINESS OPERATORS' OWN CHECKS

To decide on the classification, opening or closure of production areas, the competent authority may take into account the results of controls that food business operators or organisations representing food business operators have carried out. In that event, the competent authority must have designated the laboratory carrying out the analysis and, if necessary, sampling and analysis must have taken place in accordance with a protocol that the competent authority and the food business operators or organisation concerned have agreed.

CHAPTER III: OFFICIAL CONTROLS CONCERNING PECTINIDAE, MARINE GASTROPODS AND ECHINODERMS WHICH ARE NOT FILTER FEEDERS HARVESTED OUTSIDE CLASSIFIED PRODUCTION AREAS

Official controls on pectinidae, marine gastropods and echinoderms, which are not filter feeders, harvested outside classified production areas shall be carried out in fish auctions, dispatch centres and processing establishments.

Such official controls must verify compliance with the health standards for live bivalve molluscs laid down in Annex III, Section VII, Chapter V, of Regulation (EC) No 853/2004 as well as compliance with other requirements of Annex III, Section VII, Chapter IX, of that Regulation.
ANNEX III

FISHERY PRODUCTS

CHAPTER I: OFFICIAL CONTROLS OF PRODUCTION AND PLACING ON THE MARKET

1. Official controls on the production and placing on the market of fishery products are to include, in particular:

   (a) a regular check on the hygiene conditions of landing and first sale;

   (b) inspections at regular intervals of vessels and establishments on land, including fish auctions and wholesale markets, to check, in particular:

      (i) where appropriate, whether the conditions for approval are still fulfilled,

      (ii) whether the fishery products are handled correctly,

      (iii) for compliance with hygiene and temperature requirements,

      and

      (iv) the cleanliness of establishments, including vessels, and their facilities and equipment, and staff hygiene;

      and

   (c) checks on storage and transport conditions.

2. However, subject to paragraph 3, official controls of vessels:

   (a) may be carried out when vessels call at a port in a Member State;

   (b) concern all vessels landing fishery products at ports in the Community, irrespective of flag;

   and

   (c) may, if necessary, when the competent authority of the Member State the flag of which the vessel is flying carries out the official control, be carried out while the vessel is at sea or when it is in a port in another Member State or in a third country.

3. (a) In the case of an inspection of a factory or freezer vessel flying the flag of a Member State carried out with a view to the approval of the vessel, the competent authority of the Member State the flag of which the vessel is flying is to carry out inspections in such a manner as to comply with the requirements of Article 3, particularly the time limits of Article 3(2). If necessary, that competent authority may inspect the vessel while it is at sea or when it is in a port in another Member State or in a third country.

   (b) When the competent authority of the Member State the flag of which the vessel is flying has granted the vessel conditional approval in accordance with Article 3, that competent authority may authorise a competent authority of:

      (i) another Member State,

      or

      (ii) a third country that appears on a list of third countries from which imports of fishery products are permitted drawn up in accordance with Article 11, to carry out a follow-up inspection with a view to granting full approval or prolonging conditional approval in accordance with Article 3(1)(b) or to keeping approval under review in accordance with Article 3(4). If necessary, that competent authority may inspect the vessel while it is at sea or when it is in a port in another Member State or in a third country.
4. When the competent authority of a Member State authorises the competent authority of another Member State or of a third country to carry out inspections on its behalf in accordance with paragraph 3, the two competent authorities are to agree on the conditions governing such inspections. These conditions are to ensure, in particular, that the competent authority of the Member State the flag of which the vessel is flying receives reports on the results of inspections and on any suspected non-compliance without delay, so as to enable it to take the necessary measures.

CHAPTER II: OFFICIAL CONTROLS OF FISHERY PRODUCTS

Official controls of fishery products are to include at least the following elements.

A. ORGANOLECTIC EXAMINATIONS

Random organoleptic checks must be carried out at all stages of production, processing and distribution. One aim of these checks is to verify compliance with the freshness criteria established in accordance with Community legislation. In particular, this includes verifying, at all stages of production, processing and distribution, that fishery products at least exceed the baselines of freshness criteria established in accordance with Community legislation.

B. FRESHNESS INDICATORS

When the organoleptic examination reveals any doubt as to the freshness of the fishery products, samples may be taken and subjected to laboratory tests to determine the levels of total volatile basic nitrogen (TVB-N) and trimethylamine nitrogen (TMA-N).

The competent authority is to use the criteria laid down under Community legislation.

When the organoleptic examination gives cause to suspect the presence of other conditions which may affect human health, appropriate samples are to be taken for verification purposes.

C. HISTAMINE

Random testing for histamine is to be carried out to verify compliance with the permitted levels laid down under Community legislation.

D. RESIDUES AND CONTAMINANTS

Monitoring arrangements are to be set up to control the levels of residues and contaminants in accordance with Community legislation.

E. MICROBIOLOGICAL CHECKS

Where necessary, microbiological checks are to be performed in accordance with the relevant rules and criteria laid down under Community legislation.

F. PARASITES

Random testing is to take place to verify compliance with Community legislation on parasites.

G. POISONOUS FISHERY PRODUCTS

Checks are to take place to ensure that:

1. fishery products derived from poisonous fish of the following families are not placed on the market: Tetraodontidae, Molidae, Diodontidae and Canthigasteridae;
2. fresh, prepared, frozen and processed fishery products belonging to the family Gempylidae, in particular Ruvettus pretiosus and Lepidocybium flavobrunneum, may only be placed on the market in wrapped/packaged form and must be appropriately labelled to provide information to the consumer on preparation/cooking methods and on the risk related to the presence of substances with adverse gastrointestinal effects. The scientific names of the fishery products and the common names must appear on the label;

3. fishery products containing biotoxins such as ciguatera or other toxins dangerous to human health are not placed on the market. However, fishery products derived from bivalve molluscs, echinoderms, tunicates and marine gastropods may be placed on the market if they have been produced in accordance with Section VII of Annex III to Regulation (EC) No 853/2004 and comply with the standards laid down in Chapter V, point 2, of that Section.

CHAPTER III: DECISIONS AFTER CONTROLS

Fishery products are to be declared unfit for human consumption if:

1. organoleptic, chemical, physical or microbiological checks or checks for parasites have shown that they are not in compliance with the relevant Community legislation;

2. they contain in their edible parts contaminants or residues in excess of the limits laid down in Community legislation or at levels where the calculated dietary intake would exceed the acceptable daily or weekly intake for humans;

3. they derive from:
   (i) poisonous fish,
   (ii) fishery products not complying with the requirement of part G, point 2, of Chapter II concerning biotoxins,
      or
   (iii) bivalve molluscs, echinoderms, tunicates or marine gastropods containing marine biotoxins in total quantities exceeding the limits referred to in Regulation (EC) No 853/2004;
      or
   (iv) bivalve molluscs, echinoderms, tunicates or marine gastropods containing marine biotoxins in total quantities exceeding the limits referred to in Regulation (EC) No 853/2004;

4. the competent authority considers that they may constitute a risk to public or animal health or are for any other reason not suitable for human consumption.
CHAPTER I: CONTROL OF MILK AND COLOSTRUM PRODUCTION HOLDINGS

1. Animals on milk and colostrum production holdings must be subject to official controls to verify that the health requirements for raw milk and colostrum production, and in particular the health status of the animals and the use of veterinary medicinal products, are being complied with.

These controls may take place at the occasion of veterinary checks carried out pursuant to Community provisions on animal or public health or animal welfare and may be carried out by an approved veterinarian.

2. If there are grounds for suspecting that the animal health requirements are not being complied with, the general health status of the animals is to be checked.

3. Milk and colostrum production holdings are to undergo official controls to verify that hygiene requirements are being complied with. These official controls may involve inspections and/or the monitoring of controls that professional organisations carry out. If it is shown that the hygiene is inadequate, the competent authority is to verify that appropriate steps are taken to correct the situation.

CHAPTER II: CONTROL OF RAW MILK AND COLOSTRUM UPON COLLECTION

1. In the case of raw milk and colostrum, the competent authority is to monitor the checks carried out in accordance with Annex III, Section IX, Chapter I, Part III to Regulation (EC) No 853/2004.

2. If the food business operator has not corrected the situation within three months of first notifying the competent authority of non-compliance with the criteria with regard to plate count and/or somatic cell count, delivery of raw milk and colostrum from the production holding is to be suspended or — in accordance with a specific authorisation of, or general instructions from, the competent authority — subjected to requirements concerning its treatment and use necessary to protect public health. This suspension or these requirements are to remain in place until the food business operator has proved that the raw milk and colostrum again complies with the criteria.
ANNEX V

ESTABLISHMENTS NOT SUBJECT TO THE LISTING REQUIREMENT OF ARTICLE 12(1)

The following third-country establishments need not appear on lists drawn up and updated in accordance with Article 12(4):

1. establishments handling products of animal origin for which Annex III to Regulation (EC) No 853/2004 does not lay down requirements;
2. establishments carrying out only primary production;
3. establishments carrying out only transport operations;
4. establishments carrying out only the storage of products of animal origin not requiring temperature-controlled storage conditions.
ANNEX VI

REQUIREMENTS FOR CERTIFICATES ACCOMPANYING IMPORTS

1. The representative of the competent authority of the third country of dispatch issuing a certificate to accompany a consignment of products of animal origin destined for the Community must sign the certificate and ensure that it bears an official stamp. This requirement applies to each sheet of the certificate if it consists of more than one. In the case of factory vessels, the competent authority may authorise the captain or another ship's officer to sign the certificate.

2. Certificates must be drawn up at least in the official language or languages of the Member State of destination and those of the Member State in which the border inspection takes place, or be accompanied by a certified translation into that language or languages. However, a Member State may consent to the use of an official Community language other than its own.

3. The original version of the certificate must accompany consignments on entry into the Community.

4. Certificates must consist of:
   
   (a) a single sheet of paper;

   or

   (b) two or more pages that are part of an integrated and indivisible sheet of paper;

   or

   (c) a sequence of pages numbered so as to indicate that it is a particular page in a finite sequence (for example, 'page 2 of four pages').

5. Certificates must bear a unique identifying number. Where the certificate consists of a sequence of pages, each page must indicate this number.

6. The certificate must be issued before the consignment to which it relates leaves the control of the competent authority of the third country of dispatch.