Aims of control

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

Conditions applicable to the embryo production team

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

1) The team should be approved by the Competent Authority.
2) The team should be supervised by a team veterinarian.
3) The team veterinarian is responsible for all team operations which include the hygienic collection of ovaries and oocytes and all other procedures involved in the production of embryos intended for international movement.
4) Team personnel should be adequately trained in the techniques and principles of disease control. High standards of hygiene should be practised to preclude the introduction of infection.
5) The production team should have adequate facilities and equipment for:
   a) collecting ovaries and/or oocytes;
   b) processing of oocytes and production of embryos at a permanent or mobile laboratory;
   c) storing oocytes and/or embryos.

These facilities need not necessarily be at the same location.

6) The embryo production team should keep a record of its activities, which should be maintained for inspection by the Veterinary Authority Services for a period of at least two years after the embryos have been exported.

7) The embryo production team should be subjected to regular inspection at least once a year by an Official Veterinarian to ensure compliance with procedures for the sanitary collection and processing of oocytes and the production and storage of embryos.
Article 4.8.3.

Conditions applicable to the processing laboratories

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

Embryos may also be subjected to any required treatments such as washing and storage and quarantine in this laboratory.

Additionally:

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

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

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

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

Article 4.8.4.

Conditions applicable to donor animals

Oocytes for the in vitro production of embryos are obtained from donors basically in two different ways: individual collection or batch collection. The recommended conditions for these differ.

Individual collection usually involves the aspiration of oocytes from the ovaries of individual live animals on the farm where the animal resides, or at the laboratory. Occasionally oocytes may also be recovered from individual live donors by aspiration from surgically excised ovaries. When oocytes are recovered from individual live animals, the conditions for these donors should resemble those set out in Article 4.7.4.

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

Batch collection involves the removal of ovaries from batches of donors slaughtered at a slaughterhouse/abattoir (hereafter ‘abattoir’); these ovaries are then transported to the processing laboratory where the oocytes are recovered from the ovarian follicles by aspiration or slicing techniques. Batch collection has the disadvantage that it is usually impractical to relate the ovaries which are transported to the laboratory to the donors which were slaughtered at the abattoir. Nevertheless, it is critical to ensure that only healthy tissues are obtained and that they are removed from the donors and transported to the laboratory in a hygienic manner.

Additionally:

1) The Veterinary Authority should have knowledge of the herd(s) or flock(s) from which the donor animals have been sourced.

2) The donor animals should not originate from herds or flocks that are subject to veterinary restrictions for foot and mouth disease, rinderpest and or pest des petits ruminants, and neither should the removal of any tissue or aspiration of oocytes take place in an infected zone, or one that is subject to veterinary restrictions for those diseases.
3) In the case of oocyte recovery from live donors, post-collection surveillance of the donors and donor herd(s) or flock(s) should be conducted based on the recognised incubation periods of the diseases of concern to determine retrospectively the health status of donors.

4) In the case of oocyte recovery from batches of ovaries collected from an slaughterhouse/abattoir, the abattoir it should be officially approved and under the supervision of a veterinarian whose responsibility is to ensure that ante-mortem and post-mortem inspections of potential donor animals are carried out, and to certify them to be free of clinical or pathological signs of the diseases listed in point 2.

5) Donor animals slaughtered at an slaughterhouse/abattoir should not have been animals designated for compulsory slaughter for a notifiable disease and or should not be slaughtered at the same time as such animals donors from which ovaries and other tissues will be removed.

6) Batches of ovaries and other tissues collected from an slaughterhouse/abattoir should not be transported to the processing laboratory before confirmation has been obtained that ante- and post-mortem inspection of donors has been satisfactorily completed carried out with favourable results.

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

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

**Article 4.8.5.**

Optional tests and treatments

A supplementary approach for ensuring that in vitro produced embryos do not transmit disease is by testing various materials to confirm the absence of pathogenic organisms agents listed in point 2 of Article 4.8.4.

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

Tests may be carried out on the following materials:

1) non-viable oocytes or embryos from any stage of the in vitro production line from batches intended for export;

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

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

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

Additionally:

1) Semen used to fertilise oocytes in vitro should have been collected and processed in accordance with Chapter 4.5, and meet the health requirements and standards set out in Chapter 4.6. as appropriate to the species.

When the donor of the semen used to fertilise the oocytes is dead, and when the health status of the semen donor concerning a particular infectious disease or diseases of concern was not known at the time of semen collection, additional tests on the spare embryos may be required to verify that these infectious diseases were not transmitted.
An alternative may be to test an aliquot of semen from the same collection date.

2) Any biological product of animal origin, including co-culture cells and media constituents, used in oocyte recovery, maturation, fertilisation, culture, washing and storage should be free of living pathogenic agents. Media should be sterilised prior to use by approved methods in accordance with the IETS Manual\(^1\) and handled in such a manner as to ensure that sterility is maintained. Antibiotics should be added to all fluids and media as recommended in the IETS Manual\(^1\).

3) All equipment used to recover, handle, culture, wash, freeze and store oocytes or embryos should be new or cleaned and sterilised prior to use as recommended in the IETS Manual\(^1\).

**Article 4.8.6.**

**Risk management**

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

1) the first phase comprises the risk potential for oocyte or embryo contamination and depends on:
   a) the disease situation in the exporting country and/or zone;
   b) the health status of the herds or flocks and the donors from which the ovaries, oocytes or embryos or semen for fertilisation of oocytes are collected;
   c) the pathogenic characteristics of the specified disease pathogenic agents listed in point 2 of Article 4.8.4.;

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

3) the third phase, which is applicable to diseases listed in point 2 of Article 4.8.4. encompasses the risk reductions resulting from:
   a) post-collection surveillance of the donors and donor herds or flocks based on the recognised incubation periods of the diseases of concern to determine retrospectively the health status of the donors whilst the embryos are stored (in species where effective storage by cryopreservation is possible) in the exporting country. Post-collection surveillance of donors is not, of course, possible in the case of batch collection from an *slaughterhouse/abattoir*, although surveillance of the herds or flocks of origin may be possible;
   b) testing of oocytes, embryos, co-culture cells, media and other samples (e.g. blood) (as referred to in Article 4.8.5.) in a laboratory for presence of disease pathogenic agents.

**Article 4.8.7.**
Conditions applicable to the storage and transport of oocytes and embryos

Oocytes and in vitro produced embryos can be stored and transported fresh, chilled or frozen.

Fresh embryos may undergo culture in portable incubators during transportation and should arrive at the recipient animal within five days, in time for transfer of the mature blastocysts. Chilled embryos should be transferred within 10 days of chilling.

The Veterinary Services should have knowledge of the variety of oocyte and embryo storage systems available and should have procedures in place for the safe and timely inspection and certification of these oocytes and embryos to ensure their viability.

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

2) For frozen oocytes and embryos
   a) Sterile ampoules, vials or straws should be sealed prior to freezing or after vitrification and should be labelled according to the IETS Manual.
   b) The frozen oocytes and embryos should, if possible, depending on the species, be frozen in fresh liquid nitrogen or other cryoprotectant and then stored in fresh cryoprotectant liquid phase nitrogen or in the vapour phase of liquid nitrogen cleaned in disinfected containers under strict hygienic conditions at a storage place.
   c) Liquid nitrogen containers should be sealed prior to shipment.

Rationale: The United States recommends the word “fresh” be retained or replaced with “not previously used” – removing the term “fresh” might be misinterpreted to mean that previously used liquid nitrogen is acceptable for storage following the initial freezing. The addition of the word “in” provides clarity and is grammatically correct.

3) For fresh or chilled oocytes and embryos
   a) Sterile ampoules, vials or straws should be sealed prior to storing in portable incubators at the time of freezing and should be labelled in accordance with the IETS Manual.
   b) The fresh or chilled oocytes and embryos should be stored under strict hygienic conditions in portable incubators disinfected in accordance with the IETS Manual and manufacturer’s instructions.
   c) Portable incubators should be sealed prior to shipment.

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

4.8.7.2.b: Oocytes and embryos should not be exported until the appropriate veterinary certificates are completed.

Article 4.8.7.2.b:

b) The frozen oocytes and embryos should, if possible, depending on the species, be frozen in fresh liquid nitrogen or other cryoprotectant and then stored in fresh cryoprotectant liquid phase nitrogen or in the vapour phase of liquid nitrogen cleaned disinfected containers under strict hygienic conditions at a storage place.

Should be changed to:

b) The frozen oocytes and embryos should, if possible, depending on the species, be frozen in fresh liquid nitrogen or other cryoprotectant and then stored in fresh cryoprotectant liquid phase nitrogen or in the vapour phase of liquid nitrogen cleaned in disinfected containers under strict hygienic conditions at a storage place.
Article 4.8.8.

Procedure for micromanipulation

When micromanipulation of the embryos is to be carried out this should be done after completion of the treatment described in point 2 of Article 4.8.6. and conducted in accordance with Chapter 4.9.