Preservation and shipment of chilled and cryopreserved dog semen

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SUMMARY

The transport and artificial insemination of chilled (4°C) and cryopreserved (-196°C) dog semen have gained increasing interest worldwide and have become very popular among dog breeders. Whereas cryopreservation of dog sperm is a complicated and time consuming procedure, which is almost exclusively performed at universities, the chilling of dog semen can be handled by veterinarians in their private practices, provided that the basic knowledge of chilling and diluting semen is acquired. Immediately after sperm collection, the quality of the fresh sample is evaluated and recorded before diluting in an appropriate extender. Subsequently, the diluted semen is gradually chilled to 4°C. It can be stored at 4°C for several days or transported in a thermos flask, a styrofoam box or a Minitübe neopore box. Cryopreserved dog sperm is mostly transported in a dry-shipper container. The rules and legislation for the shipment of chilled and frozen dog semen are rather complicated. They differ between almost every country and may change over time. To comply with all the administrative procedures, it is necessary to plan the transport of semen well in advance.

INTRODUCTION

Artificial insemination (AI) in dogs can be performed with fresh, chilled (4°C) or frozen-thawed (-196°C) semen. Artificial insemination with fresh semen was first described by the Italian priest Spallanzani in 1789. However, it was not until 1954 that the first successful AI using chilled (4°C) dog semen was reported by Harrop. Fifteen years later, Seager (1969) reported the first live offspring following AI with frozen-thawed (-196°C) canine semen. Because of the obvious advantages and possibilities, AI with chilled and frozen-thawed semen gained increasing interest amongst dog breeders, veterinarians and experimental research facilities worldwide. Chilled and frozen semen makes international transport of genetic material possible, as it is easier and cheaper than the transport of animals. Based on a large study performed in Sweden, the possibility for the international exchange of genetic material is the most important reason for performing AI in dogs (Linde-Forsberg and Forsberg, 1993). Moreover, AI with chilled and especially frozen-thawed semen makes it possible to transfer valuable genetic material at a later time (even after the death of the dog) and has several economic and sanitary advantages (Rijsselaere et al., 2001, 2010; Van Soom et al., 2001). Currently, in Europe, the percentages of canine AI’s performed by veterinarians using either fresh, chilled or frozen-thawed semen are 50-55%, 10% and 35-40%, respectively (Linde-Forsberg et al., 2010).

SPERM COLLECTION AND EVALUATION

Dog sperm is generally collected by digital manipulation and stimulation of the bulbus penis (Linde-Forsberg, 1991). A quiet environment and the presence of a teaser bitch are advised for less experienced dogs. Alternatively, a vaginal swab of a bitch in oestrus which was collected previously and stored in the freezer (-20°C), can be used to stimulate the male. In case of experienced stud dogs however, sperm collection is usually possible without any extra stimulation. The three fractions of the ejaculate are collected in separate, pre-warmed plastic vials. The first and the third fractions originate from the prostate (the only secondary gland in the dog); the second sperm-rich fraction usually consists of 0.5 to 5 ml sperm.

Immediately after collection, the sperm-rich fraction is evaluated macroscopically (volume, color, admixtures and homogeneity) and microscopically (motility, concentration, morphology and membrane integrity) especially when the sample is to be chilled or frozen and transported. Motility (total and progressive) can be assessed subjectively on a pre-warmed glass slide on a scale from 0 to 100%. The sperm concentration can be determined using a counting chamber (e.g. Bürker or Thoma chamber) after a 1:40 dilution with tap water or diluted formol resulting in immobility of the spermatozoa. The percentage of normal and membrane intact spermatozoa is generally examined on eosin-nigrosin stained smears by assessing at least 100 spermatozoa individually. The morphology can also be evaluated using a Diff-Quick staining.
Concerning: Anglo of the Lembeck Farm, Border Collie, born 23/07/2001, Tattoo CHBNRC, Chip number: 52851320000075075

Owner: Mr Freddy De Boever, Kemmelstraat 75, 9100 Nieuwerkerken-Waas, Belgium

To whom it may concern,

Above mentioned Anglo of the Lembeck Farm was presented at our department on January 8th, 2011 for sperm collection, evaluation and freezing. Above mentioned Anglo of the Lembeck Farm was submitted to serological testing on January 8th, 2011 with negative result for Brucella canis and less than 50% agglutination at dilution 1:30 for Leptospira canicola and ieterohaemorrhagia.

General examination: Alerte, no abnormalities

Examination reproductive tract: Penis, prepuce, scrotum and prostate: normal
Testes and epididymides: normal - both testicles present

Evaluation of the fresh semen quality before freezing:
- Volume: 2.5 ml sperm rich fraction
- Concentration: 400 x 10⁶/ml
- Motility: 85%; Progressive motility: 80%
- Morphology: % live spermatozoa: 90%
  % normal spermatozoa: 90%

The ejaculate was frozen using the method of Rota A. et al. (1998). The total shipment contains 10 straws of approximately 100x10⁶ spermatozoa/straw of Anglo of the Lembeck Farm. Thawing should be performed immediately before artificial insemination in a warm water bath of 37°C for one minute.

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Figure 1. Example of a certificate with the results of the sperm evaluation immediately after collection.
the last decade, several new techniques for sperm assessment, such as computer assisted sperm analysis (CASA) and fluorescent staining techniques, have been described in dogs. They allow a more detailed and objective sperm assessment (Verstegen et al., 2002; Rijsselaere et al., 2005, 2007). A certificate with the results of the sperm evaluation immediately after collection should always accompany the sperm when transported. An example of a sperm certificate is shown in Figure 1.

SEMEN EXTENDERS

Since spermatozoa are metabolically relatively inert, the extracellular environment plays a crucial role in their survival. Prostatic fluid appeared not to be an appropriate medium for prolonged preservation of canine spermatozoa at 4°C (England and Allen, 1992). Spermatozoa which are preserved either chilled (4°C) or frozen (-196°C) should therefore be diluted in a proper extender. Many different extenders have been described and are currently available. They have been proven to be suitable in both in vitro and in vivo studies. In general, a semen extender should protect the spermatozoa from cold shock during cooling (by adding 10-20% egg yolk or skim milk), it should provide energy substrates (sugars such as fructose or glucose) and maintain the spermatozoa at a constant pH and osmolarity by adding buffers (England, 1993). To prevent the growth of bacteria, semen extenders should also contain some antibiotics (mostly penicillin or streptomycin). If the semen is frozen, cryoprotectants, such as glycerol, should be added. These extenders can be home-made by the veterinarian, they can be purchased at local universities or by several commercial companies. In this respect, the Tris-citric acid-egg-yolk-fructose based extenders (also referred to as the Uppsala diluents) are probably the most documented and used extenders for the preservation of dog sperm (Linde-Forsberg, 2002). Because egg yolk is a biologically hazardous compound, studies are performed replacing egg yolk by egg yolk derived phospholipids or vegetable lecithin to avoid using substances of animal origin (Farstad, 2009).

CHILLING SEMEN AND TRANSPORT

The use of chilled semen is especially interesting when the semen is to be transported over short distances (within Europe) because it is easier and cheaper than using cryopreserved semen. A good follow-up of the bitch to determine ovulation and consequently to determine the ideal insemination days, is crucial because it is better to inseminate the semen as quick as possible after collection. In general, semen can be delivered in most of the European countries within 24 to 48 hours.

To chill a sperm sample, the second sperm-rich fraction can be diluted with Tris-citric acid-fructose diluent with 20% egg yolk in a proportion of 1:3 to 1:4 (1 mL of semen with 3 to 4 mL of diluent). Subsequently, the diluted sperm sample is placed in a water bowl (at 37°C) which can subsequently be placed in the fridge and slowly be cooled (in a ‘bain-marie’) to 4 or 5°C.

Figure 2. A diluted dog sperm sample is placed in a water bowl (at 37°C) which can subsequently be placed in the fridge and slowly be cooled (in a ‘bain-marie’) to 4 or 5°C.
FREEZING SEMEN AND TRANSPORT

The use of frozen dog sperm is especially indicated when transport over larger distances is required or when the genetic material of the stud dog will be used at a later time. The freezing of dog sperm is a rather complicated and time-consuming procedure which is mostly performed at universities or large veterinary clinics. In Belgium, the kennel club Sint-Hubertus only allows dog sperm to be frozen in an official university centre (Faculty of Veterinary Medicine in Ghent or Liège) and only after approval of the stud dog by the kennel club.

Numerous procedures have been described for canine cryopreservation with variable results. Our department consistently uses the ‘Uppsala method’ with slight modifications (Linde-Forsberg 2001, 2010). In short, the second sperm-rich fraction is diluted with a Tris-citric acid-egg-yolk-fructose based extender with glycerol (3%) and chilled to 4-5°C for 1-2h (= equilibration period). Subsequently, a second extender with Equex and a higher percentage of glycerol (7%) is added and the sperm is packaged in straws (0.5 or 0.25 ml) or in pellets. Both are considered to be equally good but 0.5 ml straws are more practical to handle and label. Straws should consistently be labelled with date, name of the dog, breed, tattoo or chip number and freezing centre. Finally, the straws are placed on a rack which is located several centimetres above the level of liquid nitrogen. By making variations in the distance of the sperm straws to the level of liquid nitrogen, variable periods of time, different freezing rates are created. The above mentioned freezing technique is a ‘static method’, it is easy to conduct but allows only little control over the actual freezing rate. The repeatability of a freezing protocol can be improved when programmable automated freezing devices (= dynamic method) are used, such as the Mini Digitcool, the Planer 10 TM or IceCube (Sy-Lab) (Schafer-Somi et al., 2006), which may be useful when a large number of straws are frozen simultaneously (Figure 4). However, automated programmable freezers are very expensive. Following the cryopreservation procedure, the straws can be stored in a liquid nitrogen container (Figure 5A), if necessary for years until they are thawed and used for the insemination of a bitch or they can be transported abroad in a dry shipper (Figure 5B). The costs for the transport of dry-shippers are in general high due to the weight of the container (10-20 kg). Moreover, many airlines and transport companies are reluctant to ship them. Recently, a new single-use dry-shipper 3L (ST Technologies, USA) has been introduced. This shipper only weighs 4.5 kg when filled. It has a holding time of approximately four days and it is intended for a one-way shipment reducing the costs to about 1/4th compared to normal dry shippers (Linde-Forsberg, 2010). Additionally, on some airlines, this new dry shipper is accepted as hand luggage (Linde-Forsberg, 2010).

Before inseminating the frozen semen, the straws should be thawed in a water bath at 37°C for 30 or 60 seconds or in a water bath at 70°C for 6-7 seconds.
this respect, it is very important to consistently follow the thawing instructions of the freezing centre which should always accompany the shipment!

FREEZING OF CHILLED SEMEN

Several studies have shown that semen which was previously chilled for two to three days at 4°C, can subsequently be frozen successfully. This makes it possible to collect, to chill and transport the dog semen to a nearby semen bank where it can be cryopreserved and stored at -196°C for AI at a later time (Verstegen et al., 2005; Hermansson and Linde-Forsberg, 2006).

REGULATIONS AND RECOMMENDATIONS FOR SHIPMENT

The rules for the shipment of chilled and frozen dog semen are rather complicated and differ from country to country. Some countries such as Belgium are ‘easy’ (category 1), while other countries such as Australia, are very strict regarding the import of dog semen (category 4). Original import permits, a health certificate, several serological tests (for Brucella Canis and Leptospirosis) are only some of the requirements. The importation of chilled and cryopreserved dog semen into Belgium however, is currently not subject to any restrictions, due to the lack of regulations. It should also be kept in mind that national legislation and regulations applied by the local kennel club may change in time. Moreover, the rules for importing chilled semen may be different from those applying to frozen semen. An excellent overview of the regulations and recommendations for shipment of chilled and frozen dog semen including many countries worldwide is published by Linde-Forsberg (2001) on the IVIS website (http://www.ivis.org/home.asp). It is therefore very important to advise the importer or dog owner to contact the Ministry of Agriculture (or local authority) to inform them about the current rules and regulations of a particular country. This should be done well in advance of the planned sperm collection and transport in order to be able to carry out all the necessary blood tests, health requirements and paper work.

CONCLUSION

The choice whether to use chilled or frozen dog semen for transport depends on many aspects, such as the distance (chilled semen is mostly used for transport within a country or within Europe whereas overseas shipments are frequently performed using cryopreserved semen), the purpose of the semen (AI of only one bitch or AI of several bitches), the cost (cryopreserved sperm transport is often more expensive regarding the cryopreservation procedure and the shipping costs of the dry shippers), the local regulations (may differ for chilled or cryopreserved semen), the personal preference of the owner, the know-how and skills for inseminating chilled and frozen semen of the responsible veterinarian, etc. An overview with the different insemination techniques and the results which can be obtained following AI in dogs has recently been published by Rijsselaere et al. (2010).

REFERENCES


