

OVERVIEW OF TURKEY SEMEN STORAGE: FOCUS ON CRYOPRESERVATION – A REVIEW

Nicolaia Iaffaldano1+, Michele Di Iorio1, Silvia Cerolini2, Angelo Manchisi1

¹Department of Agricultural, Environmental and Food Sciences, University of Molise, via De Sanctis snc, 86100 Campobasso, Italy ²Department of Health, Animal Production and Food Safety, University of Milan, via Trentacoste 2, 20134 Milan, Italy *Corresponding author: nicolaia@unimol.it

Abstract

This review updates the current state of technologies available for turkey semen storage (hypothermic-liquid storage and cryopreservation), with special attention paid to cryopreservation. Liquid semen can be stored for up to 24 or 48 h at temperatures around 5°C, while cryopreservation allows long-term storage at -196°C. The possibility of using frozen turkey semen for artificial insemination (AI) would have practical benefits for turkey production. Reported fertility rates in response to AI using frozen/thawed semen range from 15.8 to 84.3%. Unsatisfactory fertility may be attributed to an inability of turkey spermatozoa to successfully survive the freezing/thawing process, and this, along with the high variability observed, makes this technique unacceptable for commercial breeding programs. There is therefore a need to standardize the whole freezing and thawing process to improve the post-thaw quality of turkey semen and minimize variability in results. Finding an efficient freezing protocol for turkey semen will allow for the creation of a sperm cryobank, improving current prospects for the commercial use of frozen turkey semen and also for the long-term conservation of the genetic diversity of this bird.

Key words: turkey semen, cryopreservation, semen storage, semen quality, fertility

The cryopreservation and storage of germplasm has long been valued for the indefinite preservation of genetic material, especially in cases of high-risk populations. An immediate need for this practice was identified for research using unique poultry lines (Long and Kulkarni, 2004). Today, however, semen cryopreservation seems to be the only effective method of storing reproductive cells for the *ex situ* management of genetic diversity in birds (Blesbois, 2011; Kowalczyk and Łukaszewicz, 2015). Successful semen cryopreservation has enabled the creation of semen banks for several wild and some poultry species (Saint Jalme et al., 2003; Blackburn, 2006; Woelders et al., 2006; Blesbois, 2007; Blanco et al., 2009; Kowalczyk et al., 2012). However, research efforts have not yet served to create a turkey semen cryobank. The possibility of using turkey semen in frozen form for artificial insemination (AI), besides maintaining and ensuring the long-term conservation of this bird's genetic diversity, would have practical benefits for turkey production. Turkeys are the only commercial poultry species that depend entirely upon AI for fertile egg production. This is because the difference in size between males and females of commercial strains, resulting from genetic selection, makes it impossible for turkeys to naturally mate (Donoghue and Wishart, 2000; Iaffaldano et al., 2010). When freshly collected semen is used, AI yields exceptional fertility rates. However, this practice requires that toms and hens are kept on the same farm, meaning that a large number of birds have to be managed (Long and Bakst, 2007; Rosato et al., 2012). The turkey industry would therefore greatly benefit if semen could be stored after collection and used for subsequent AI (Rosato et al., 2012). The technologies available for semen storage are essentially: 1) hypothermic-liquid storage (refrigeration), which enables the storage of semen for up to 24 or 48 h at chilling temperatures around 5°C (Wishart, 2009); and 2) cryopreservation, whereby semen can be long-term stored at -196° C, the temperature of liquid nitrogen. In this review, we update the current state of these technologies, with special attention paid to the cryopreservation of turkey semen.

Hypothermic-liquid storage

During liquid storage at reduced temperature, sperm metabolism is not completely arrested. Using this method, domestic turkey semen cannot be stored longer than 6 h without losing its fertilizing ability, even when oxygenated and stored with appropriate diluents (Thurston, 1995). Recently, the research focus has been on defining the optimum diluent and improving storage systems addressing the composition of seminal plasma and sperm metabolic requirements or through the use of antioxidant supplements with biotechnologies designed to optimize the quality of stored turkey semen and consequent fertility (Donoghue and Wishart, 2000; Douard et al., 2000; Neuman et al., 2002; Douard et al., 2003; Iaffaldano and Meluzzi, 2003; Long and Kramer, 2003; Douard et al., 2004, 2005; Iaffaldano et al., 2005; Dimitrov et al., 2007; Iaffaldano et al., 2008; Zaniboni and Cerolini, 2009; Rosato et al., 2012). The results of these studies, although variable, have indicated the compromised survival of chilled turkey semen determining a reduction both in its quality and fertilizing ability (Table 1). The most popular extender used for stored turkey semen is BPSE (Beltsville Poultry Semen Extender). Extenders are buffers that promote the immediate survival of spermatozoa because they provide a similar osmotic pressure (330-400 mOsm) and pH (7.0-7.5) to that of seminal plasma, and are also a source of energy due to substrates such as carbohydrates (glucose or fructose) or other components such as citrate, glutamate and acetate (Iaffaldano et al., 2005). The liquid refrigeration of semen has returned better results in terms of both quality and fertilizing ability in the chicken (Sexton and Fewlass, 1978; Blesbois et al., 1999; de Figueiredo et al., 1999; Lemoine et al., 2011) and duck (Kasai et al., 2000; Penfold et al., 2001) (Table 1). Accordingly, Donoghue and Wishart (2000) reported fertility levels comparable to inseminated fresh semen for the storage at refrigeration temperatures of chicken semen for up to 24 h and of turkey semen for up to 6 h only. Moreover, turkey spermatozoa are active only in aerobic conditions whereas chicken spermatozoa are active also in anaerobic conditions. Turkey spermatozoa are considered efficient because of their high oxidation rate and low lactic acid accumulation in the presence of oxygen (Sexton, 1974). Further factors affecting the storability of turkey semen are age and strain, which were found to impact the quality of both fresh semen and of semen stored in liquid form (Iaffaldano et al., 2008).

Species	Extenders	Storage time (h)	Viability (%)	Motility (%)	Fertility (%)	Reference
Turkey	BPSE	48	95	56	_	Douard et. al., 2000
	BPSE	48	35-80	40–75	_	Iaffaldano et al., 2005, 2008; Rosato et al., 2012
	BPSE	48	83–95	20-50	53-76 (24 h)	Douard et al., 2005
	CE*	48	83	35	_	Zaniboni et al., 2009
	BPSE	24	50	20-30	45-50	Long and Conn, 2012
	CE*	48	_	70–75	_	Słowińska et al., 2013
Chicken	BPSE	24	-	_	50-60	Sexton and Fewlass, 1978
	BPSE	48	93	53	_	Blesbois et al., 1999
	BPSE/Lake/ CE*	24	_	-	87–95	De Figueiredo et al., 1999
	BPSE	48	84	90	_	Lemoine et al., 2011
Duck	BPSE	24	_	85	52	Kasai et al., 2000
	BPSE	48 72	98 98	84 77	80 43	Penfold et al., 2001

Table 1. Recovered viability, motility and fertility rates recorded for chilled turkey semen (T: 4–5°C storage time: 24–72 hours) compared to rates reported for other poultry species

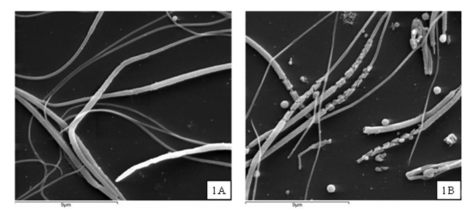
*CE: commercial extender.

Cryopreservation

The possibility of using semen in frozen form for AI is a key factor for ensuring the long-term conservation of genetic diversity through the creation of a semen cryobank. Frozen semen also has several practical advantages for turkey production. Over 50 years ago, the discovery of glycerol's cryoprotective properties led to the development of a technology for the cryopreservation of semen in a wide range of species (Long, 2006). However, despite the good progress made in the cryopreservation of semen in cattle, this preservation method has not been as successful in avian species. The reason for this is the high cost of preparing and storing frozen ejaculates compared to the market price of day-old chicks, and also low semen quality and consequently the fertility levels achievable with frozen/thawed spermatozoa (Blesbois, 2007; Iaffaldano et al., 2011). The poor fertilization rates obtained for avian as opposed to mammalian species are attributable to the unique morphological characteristics of avian spermatozoa, such as their filiform shape, long tail and condensed nucleus, which makes them more susceptible to freezing damage (Donoghue and Wishart, 2000; Long, 2006). In addition, membrane damage induced by cryopreservation results in impaired sperm transport and survival in the female reproductive tract with the consequent decreased duration of fertility that has been correlated with the number of spermatozoa in sperm storage tubules at the utero-vaginal junction (Pierson et al., 1988; Tajima, 2013).

Despite this, a variety of semen cryopreservation protocols involving different cryoprotective agents (CPAs), packaging methods, and freezing and thawing rates, have been developed, firstly in the chicken and then in other domesticated birds, such as turkey, duck and goose (see reviews Lake, 1986; Bellagamba et al., 1993; Hammerstedt, 1995; Surai and Wishart, 1996; Donoghue and Wishart, 2000; Blesbois, 2007, 2011).

Semen cryopreservation involves several steps, each one affecting sperm structure and function (Garner et al., 1999; Bailey et al., 2003): extension, cooling, CPA addition, freezing, and thawing (Bailey et al., 2003). Deleterious effects are the result of osmotic stress, and temperature changes produced during cooling, freezing and rewarming, ice crystal formation being one of the main biophysical mechanisms of sperm death (Swain and Smith, 2010) (Figure 1). A principal challenge for the survival of cells during cryopreservation, is the lethality of the intermediate temperature zone (-15 to -60° C), which is crossed twice during the cryogenic cycle, as cells are cooled and rewarmed (Gao and Critser, 2000; Blanco et al., 2011). Thus, the cryopreservation process causes numerous negative effects including damage to cell membranes (plasma and mitochondrial) and, in some cases, to the nucleus with devastating consequences for sperm survival (Blesbois, 2007). Following cryopreservation, metabolic damage may affect levels of adenosine triphosphate (ATP), which is essential for sperm motility (Long, 2006).



Bars represent 9 μ m. (1A) Fresh semen: spermatozoa show intact membranes. (1B) Cryopreserved semen: spermatozoa show damaged membranes. Scanning electron micrographs from the Department of Agricultural, Environmental and Food Sciences, University of Molise.

Figure 1. Pictures of fresh and frozen turkey semen, obtained by SEM (scanning electron microscopy)

Despite their similar morphology, the cryosurvival of sperm cells also varies among different avian species and has been correlated with the freezing procedure (i.e. speed of freezing/warming, sample volume, CPA, and CPA concentration) (Massip et al., 2004).

The decline in spermatozoa quality following semen storage is accompanied by changes in the proacrosin/acrosin system, which is involved in the acrosome reaction of spermatozoa and in phospholipids (Douard et al., 2000, 2004; Kotłowska et al., 2007; Słowińska et al., 2012, 2013). Thus, disruption of the proacrosin/acrosin system in turkey spermatozoa has been linked to a decline in semen quality during storage (Kotłowska et al., 2007; Słowińska et al., 2007; Słowińska et al., 2017; Słowińska et al., 2012). In particular, the phospholipid profile and contents of turkey spermatozoa are severely affected by *in vitro* storage, and changes in phospholipids are paralleled by the decrease in semen quality. This effect could be the consequence of endogenous metabolism of membrane phospholipid fatty acids inducing membrane destabilization. However, we cannot rule out a combination of many complex factors, including phospholipid lysis, endogenous metabolism, and lipid peroxidation (Douard et al., 2000, 2004; Kotłowska et al., 2007).

There is currently scarce information in the literature on turkey semen cryopreservation, since the freezing and thawing procedures developed for chickens or other birds are inefficient for turkey spermatozoa. Research efforts have focused on developing freezing protocols for the improved cryopreservation of turkey semen reducing the cell damage caused by freezing and thawing. Studies to date have examined the effectiveness of several cryoprotectants (glycerol, dimethylsulfoxide (DMSO), ethylene glycol, dimethylacetamide (DMA)), rapid or slow freezing-thawing procedures and the use of pellets or straws for packaging. The choice of CPA and its concentration certainly seems among the most important factors for an effective turkey semen freezing protocol. Permeable CPAs penetrate the sperm cell, increase membrane fluidity and partially dehydrate the cell, lowering the freezing point and thus reducing the formation of intracellular ice crystals, which is to be avoided because it causes physical and chemical stress. The permeable CPAs mainly used in freezing protocols for turkey semen are: glycerol, DMSO and DMA at different concentrations (Table 2). DMA and DMSO have been used as alternatives to glycerol since its discovered contraceptive effect. Glycerol has to be removed before AI which is an important drawback (Hammerstedt and Graham, 1992). Early studies examined the use of DMSO concentrations of 4% (Bakst and Sexton, 1979; Sexton, 1981). However, Iaffaldano et al. (2016) reported that 10% DMSO was better than 4%. These authors also showed (Iaffaldano et al., 2011) that a DMA concentration of 8% worked better than one of 6% when cells were frozen at high cooling rates, for example when directly plunging semen droplets into liquid nitrogen. Blanco et al. (2011) reported that 10% and 18% DMA provided more cryoprotection for turkey sperm when cryovials were used as the packing system. In this regard the effectiveness of sperm cryopreservation may also depend on the interaction between the cryoprotectant type used and the semen freezing and packaging method employed, that is the use of pellets or straws (Tselutin et al., 1999; Abouelezz et al., 2015). In addition, permeable CPAs themselves could paradoxically have a toxic effect on sperm, causing membrane destabilization and protein and enzyme denaturation; this toxicity is directly related to the CPA concentration used and the time of cell exposure (Swain and Smith, 2010).

	-		-
CPA	Freezing methods	Packaging system	Reference
DMA	– rapid	- pellets (50-100 µL)	Tselutin et al., 1995; Blanco et
(4–26%)	- slow (programmable freezer	- straws (0.25-0.5 mL)	al., 2000, 2011, 2012; Labbé et
	and liquid nitrogen vapor)	- cryovials (100-500 μL)	al., 2003; Blesbois et al., 2005;
			Iaffaldano et al., 2009, 2011, in
			press; Long et al., 2014
DMSO	- slow (programmable freezer	- straws 0.25 mL	Bakst and Sexton, 1979; Sexton,
(4–10%)	and liquid nitrogen vapor)	- cryovials	1981; Iaffaldano et al., 2016
Glycerol	- slow (programmable freezer	- straws 0.25 mL	Pandian et al., 2011; Long et al.,
(9–11%)	and liquid nitrogen vapor)		2014

Table 2. Freezing methods and devices used for the cryopreservation of turkey semen

The addition of non-permeable CPAs to the freezing medium therefore serves to offset the cryodamage caused by permeating CPAs. At similar concentrations, these substances are less toxic than permeable CPAs and have multiple protective roles such as inhibiting ice crystal growth and helping the sperm to stabilize internal solute concentrations under osmotic stress. This reduces the amount of penetrating CPAs needed (Swain and Smith, 2010). Non-penetrating cryoprotectants are generally large molecules such as polymers, sugars, proteins or amino acids (Blanco et al., 2011; Rosato and Iaffaldano, 2013; Iaffaldano et al., 2014).

Blanco et al. (2011) tested trehalose and/or sucrose as non permeable CPAs in combination with DMA and reported the improved post-thawing motility of turkey semen compared with the use of DMA alone as CPA, which was dependent upon DMA concentration.

With regard to freezing rates, two main sperm cryopreservation techniques have been tested: slow freezing (conventional freezing) and ultra-rapid freezing (Table 2). The liquid nitrogen vapors used in conventional freezing produce a step-wise decrease in temperature, whereas ultra-rapid freezing rapidly solidifies the semen sample avoiding ice crystal formation.

As packaging systems pellets, cryovials or straws have been used (Table 2) (Sexton, 1981; Zavos and Graham, 1983; Lake, 1986; Tselutin et al., 1995; Blesbois et al., 2005; Blanco et al., 2011, 2012; Iaffaldano et al., 2011; Long et al., 2014). The semen thawing procedure (temperature and time) is also crucial for the post-freezing quality of semen. During slow thawing (low temperature, long time), the small ice crystals formed during freezing start to melt, turning into large crystals (recrystal-lization) that are harmful to the spermatozoa (Watson, 1995). During fast thawing (high temperature, short time) the time for recrystallization to occur is limited and this increases the survivability of spermatozoa. Using the pellet procedure, Iaffaldano et al. (2011) observed that thawing at a temperature of 75°C for 10 sec was better than 60°C for 12 sec, while when straws were used, thawing conditions of 50°C/10 sec were more efficient than 4°C/5 min (Iaffaldano et al., 2016).

Semen cryopreservation by the pellet procedure

Among the freezing systems assessed for the cryopreservation of avian spermatozoa, the pellet procedure is cheap, easily adaptable to field conditions, takes only a few seconds for cooling and warming, the cryoprotectant does not have to be removed and very fast freezing rates are achieved compared to the glycerol/slow-freeze method (Tselutin et al., 1999). The pellet procedure consists of the use of DMA and rapid cooling through direct plunging of semen droplets into liquid nitrogen to form frozen pellets. The method was developed by Tselutin et al. (1995) and subsequently adopted by others (Surai and Wishart, 1996; Blesbois et al., 2005; Iaffaldano et al., 2009). Recently, by testing different critical step combinations, we identified the best pellet procedure as: Tselutin extender, dilution rate 1:4, semen cooling 60 min; DMA 8%; equilibration time 5 min, drop volume 80 µL and thawing at 75°C for 10–12 sec (Iaffaldano et al., 2011).

This best pellet cryopreservation protocol returned recovery rates of 60% for mobility, 40% for viability (evaluated using the LIVE/DEAD Sperm Viability Kit, by fluorescence probes SYBR-14 and PI) and 42% for osmotic tolerance (using hypoosmotic swelling test). In in vitro studies, similar (Blesbois et al., 2005; Lemoine et al., 2011) or lower (Słowińska et al., 2012) sperm quality were observed after pellet cryopreservation following different turkey semen processing conditions. In vivo, Tselutin et al. (1995) obtained encouraging results using turkey semen cryopreserved by the pellet method. Rates of fertile eggs ranged from 71 to 84.3% but, unfortunately, the semen processing conditions and in vitro sperm quality were not specified. In later studies, however, lower fertility rates of 35-38% were observed for the pellet procedure (Labbé, 2003). The good fertility results obtained by Tselutin et al. (1995) were also confirmed in chicken and ducks (Tselutin et al., 1995). It is normally recognized that turkey sperm is much more sensitive to cryodamage than chicken sperm, and that the female reproductive tract is more stringently selective of turkey spermatozoa (Blesbois, 2007; Blanco et al., 2000; Whishart, 2009). In the review by Blesbois (2007), success rates for freezing/thawing of chicken and gander semen were higher compared to the semen of most other domestic birds including the turkey. In a prior work we observed the similar susceptibility to cryopreservation of chicken and turkey semen. Recovery rates of viable sperm were 39% and 41% respectively, while pheasant semen returned a recovery rate of viable sperm of only 20% (Cerolini et al., 2009).

Fast cooling by directly plunging of cryovials into liquid nitrogen has been found to be more detrimental for turkey sperm viability than slow cooling, whereas the reverse is true for chicken semen (Blanco et al., 2000).

Variation in the cryotolerance of male gametes among different bird species is thought to be a consequence of the bird's lipid profile, including cell membrane lipids such as cholesterol and phospholipids, whose ratio determines membrane fluidity (Blesbois et al., 2005; Blanco et al., 2000; 2008); any biochemical changes occurring during cryostorage (Long, 2006); and osmotic tolerance, which is low for turkey spermatozoa (Blanco et al., 2008). Thus, there is strong evidence to suggest that the success of a semen freezing procedure in one bird species may not translate to its success in another species (reviewed in Blesbois and Brillard, 2007; Blesbois, 2011, 2012; Tajima, 2013).

			1		
Species	Packaging system	CPA concentration	Viability (%)	Motility (%)	Reference
Turkey	pellet	DMA 6%	38	54	Lemoine et al., 2011
	I		_	24	Słowińska et al., 2012
		DMA 8%	39	_	Blesbois et al., 2005
			41	59	Cerolini et al., 2009
		DMA 6-8%	30-40	55-60	Iaffaldano et al., 2009, 2011
	straws	DMSO 10%	53	48	Iaffaldano et al., 2016
		Glycerol 11%	72-84	14.8-27.5	Long et al., 2014
		DMA 6%	50.5-61.2	12.4-19.5	
	Eppendorf	DMA 18%	35-39	40	Blanco et al., 2011, 2012
Chicken	pellet	DMA 6%	33-42.5	33-50	Tselutin et al., 1999; Blesbois et al. 2005; Cerolini et al., 2009; Mocé e al., 2010; Gliozzi et al., 2011; Za niboni et al., 2014; Kowalczyk and Łukaszewicz, 2015
		DMA 3%	23.6	24.7	Abouelezz et al., 2015
		DMA 6%	18	19.8	
		Glycerol 11%	18.6	24	
	straws	Glycerol 11%	52	45	Seigneurin and Blesbois, 1995
		Glycerol 11%	79 74	-	Tselutin et al., 1999
		DMA 6% Glycerol 11%	74 54	_	Chalab et al. 1000
		DMA 6%	54 47	_	Chalah et al., 1999
		Glycerol 11%	-	50	Mocé et al., 2010
		Glycerol 11%	73	_	Peláez et al., 2011
		DMSO 11%	28	-	
		DMA 6%	60.7	46.3	Kowalczyk and Łukaszewicz, 2015
		DMA 6%	21.1	21.3	Abouelezz et al., 2015
		Glycerol 8%	37	39	
D 1	F 1.6	Glycerol 11%	47.6	43.4	
Duck	Eppendorf	DMSO 10%	-	73 68	Han et al., 2005
		Glycerol 8% DMA 10%	_	61	
		DMF 8%	_	58	
Guinea fowl	pellet	DMA 6%	31.4	-	Váradi et al., 2013
10 111	cryovials	EG 10%	41.1	_	Váradi et al., 2013
		DMF 6%	27.1	_	·
	straws	DMF 6%	19	_	Blesbois et al., 2005
DI (pellet	DMA 6%	20	17	Cerolini et al., 2009

Table 3. Effectiveness of the cryopreservation of turkey semen compared to the semen of other poultry species

Semen cryopreservation using the straw as packaging system

Although it was initially considered (Blesbois and Grasseau, 2002; Labbé et al., 2003) that straws were less efficient than pellets to restore the fertility of frozen turkey or chicken semen (Table 4), straws as a packaging system have benefits such as sperm traceability, and the safe transport of semen for breeding or storage in gene

banks. Straws have been widely used to freeze chicken semen using different CPAs such as glycerol, DMA or DMSO (Sexton, 1981; Williamson et al., 1981; Seigneurin and Blesbois 1995; Tselutin et al., 1999; Purdy et al., 2009; Mocé et al., 2010; Peláez et al., 2011; Santiago-Moreno et al., 2011). In a recent study in turkey, we obtained a semen quality recovery rate of about 50% using straws and 10% DMSO (Iaffaldano et al., 2016) (Table 3). Lower semen quality values were reported by Blanco (2011: 2012) using Eppendorf tubes as the packaging system (Table 3). In an *in vivo* study, Labbé et al. (2003) recorded lower fertility rates using the straw (15–20%) compared to the pellet procedure (Table 4). Long et al. (2014) reported similar fertility results ranging from 15.8 to 25% depending on the turkey line considered although these fertility results were achieved with just a single insemination of 240×10⁶ sperm per hen. Higher fertility has been observed for the chicken (Seigneurin and Blesbois, 1995; Tselutin et al., 1999; Chalah et al., 1999), duck (Han et al., 2005) and guinea fowl (Seigneurin et al., 2013; Váradi et al., 2013). Post-thaw quality and fertility in the chicken tends to be lower using DMSO or DMA compared with the glycerol method (Tselutin et al., 1999; Santiago-Moreno et al., 2011; Peláez et al., 2011). It is not necessary to remove these cryoprotective agents prior to artificial insemination (Tajima et al., 1990; Tajima, 2013). Glycerol concentrations before insemination, on the other hand, should be below 0.163 M to avoid its contraceptive effect as reviewed in Tajima (2013).

Species	Packaging system	CPA concentration	Fertility (%)	Reference
Turkey	pellet	DMA 4%	71-84.3	Tselutin et al., 1995
		DMA 8%	35–38	Labbé et at., 2003
	straws	DMA 8%	15-20	Labbé et al., 2003
		DMA 6%	15.8-25	Long et al., 2014
Chicken	pellet	DMA 6%	93-94.4	Tselutin et al., 1995
		DMA 6%	84.7-92.7	Tselutin et al., 1999
		DMA 3%	25	Abouelezz et al., 2015
		DMA 6%	12.8	
		Glycerol 11%	4.2	
	straws	Glycerol 11%	88	Seigneurin and Blesbois, 1995
		DMA 6%	26.7	Tselutin et al., 1999
		Glycerol 11%	63.9	
		Glycerol 11%	76	Chalah et al., 1999
		DMA 6%	88	
		DMF 6.5%	79	
		DMA 6%	84.4	Kowalczyk and Łukaszewicz, 2015
		DMA 6%	10.8	Abouelezz et al., 2015
		Glycerol 8%	28.8	
		Glycerol 11%	2.1	
Duck	flasks	DMA 5%	75.1-83.6	Tselutin et al., 1995
	Eppendorf	DMSO 10%	40	Han et al., 2005
Guinea fowl	pellet	DMA 6%	63.6	Váradi et al., 2013
	cryovials	EG 10%	29.1	Váradi et al., 2013
	straws	DMF 6%	70.7	Seigneurin et al., 2013

Table 4. Fertility rates obtained in different avian species using frozen semen

Conclusions

In conclusion, the data reviewed indicate that the commercial use of stored turkey semen, particularly in frozen form, is still not satisfactory because of the inability of turkey spermatozoa to successfully survive freezing/thawing. According to the literature, variability in the biological material used and the multiplicity of preservation procedures has meant that it has not been possible to reproduce either the quality or fertilizing capacity of stored semen. Susceptibility to semen cryopreservation varies among poultry species (Blesbois, 2007, 2011; Iaffaldano et al., 2011), within species (Siudzińska and Łukaszewicz, 2008) and/or genetic lines (Long et al., 2010, 2014), and within breeds. Though surprising, occasionally higher fertility rates obtained for frozen-thawed turkey semen may be attributed to a higher dose of spermatozoa and a greater frequency of AI (Blesbois et al., 2008). Thus, there is a clear need to standardize the whole freezing and thawing process to minimize variability in results. In addition, we need to identify the key factors in turkey semen processing that will significantly affect the success of cryopreservation. Finding an efficient freezing protocol for turkey semen and determining the appropriate inseminating dose and frequency will allow for the introduction of a sperm cryobank and improve current prospects for the commercial use of frozen turkey semen.

Acknowledgements

The authors thank Ana Burton for editorial assistance.

References

- Abouelezz F.M.K., Castaño C., Toledano-Díaz A., Esteso M.C., López-Sebastián A., Campo J.L., Santiago-Moreno J. (2015). Effect of the interaction between cryoprotectant concentration and cryopreservation method on frozen/thawed chicken sperm variables. Reprod. Dom. Anim., 50: 135–141.
- Bailey J.L., Morrier A., Cormier N. (2003). Semen cryopreservation: success and persistent problems in farm species. Canadian J. Anim. Sci., 83: 393–401.
- B a k s t M.R., S e x t o n T.J. (1979). Fertilizing capacity and ultrastructure of fowl and turkey spermatozoa before and after freezing. J. Reprod. Fert., 55: 1–7.
- Bellagamba F., Cerolini S., Cavalchini L.G. (1993). Cryopreservation for poultry semen: a review. World's Poultry Sci. J., 49: 158–166.
- Blackburn H.D. (2006). The National Animal Germplasm Program: Challenges and opportunities for poultry genetic resources. Poultry Sci., 85: 210–215.
- Blanco J.M., Gee G., Wildt D.E., Donoghue L. (2000). Species variation in osmotic cryoprotectant, and cooling rate tolerance in poultry, eagle and peregrine falcon spermatozoa. Biol. Reprod., 63: 1164–1171.
- Blanco J.M., Long J.A., Gee G., Donoghue A.M., Wildt D.E. (2008). Osmotic tolerance of avian spermatozoa: Influence of time, temperature, cryoprotectant and membrane ion pump function on sperm viability. Cryobiology, 56: 8–14.
- Blanco J.M., Wildt D.E., Hofle U., Voelker W., Donoghue A.M. (2009). Implementing artificial insemination as an effective tool for *ex situ* conservation of endangered avian species. Theriogenology, 71: 200–213.
- Blanco J.M., Long J.A., Gee G., Wildt D.E., Donoghue A.M. (2011). Comparative cryopreservation of avian spermatozoa: benefits of non-permeating osmoprotectants and ATP on turkey and crane sperm cryosurvival. Anim. Reprod. Sci., 123: 242–248.

- Blanco J.M., Long J.A., Gee G., Wildt D.E., Donoghue A.M. (2012). Comparative cryopreservation of avian spermatozoa: effects of freezing and thawing rates on turkey and sandhill crane sperm cryosurvival. Anim. Reprod. Sci., 131: 1–8.
- Blesbois E. (2007). Current status in avian semen cryopreservation. World's Poult. Sci. J., 63: 213–222.
- Blesbois E. (2011). Freezing avian semen. Avian. Biol. Res., 4: 52.
- Blesbois E. (2012). Biological features of the avian male gamete and their application to biotechnology of conservation. J. Poultry Sci., 49: 41–149.
- Blesbois E., Grasseau I. (2002). Seminal plasma affects liquid storage and cryopreservation of turkey sperm. Proc. 38th Meeting of the Society for Cryobiology, Edinburgh, UK, 29.07–01.08.2002, p. 103.
- Blesbois E., Brillard J.P. (2007). Specific features of *in vivo* and *in vitro* sperm storage in birds. Animal, 1: 1472–1481.
- Blesbois E., Grasseau I., Hermier D. (1999). Effect of vitamin E on fowl spermatozoa after liquid storage at 2 to 5 degrees C. Theriogenology, 52: 325–334.
- Blesbois E., Grasseau I., Seigneurin F. (2005). Membrane fluidity and the ability to survive cryopreservation in domestic bird spermatozoa. Reproduction, 129: 371–378.
- Blesbois E., Grasseau I., Seigneurin F., Mignon-Grasteau S., Saint Jalme M., Mialon-Richard M.M. (2008). Predictors of semen cryopreservation in chickens. Theriogenology, 69: 252–261.
- Cerolini S., Zaniboni L., Mangiagalli M.G., Cassinelli C., Marzoni M., Castillo A., Romboli I., Rosato M.P., Iaffaldano N. (2009). Sperm cryopreservation by the pellet method in chickens, turkeys and pheasants: a comparative study. Avian. Biol. Res., 1: 1758–1559.
- Chalah T., Seigneurin F., Blesbois E., Brillard J.P. (1999). *In vitro* comparison of fowl sperm viability in ejaculates frozen by three different techniques and relationship with subsequent fertility *in vivo*. Cryobiology, 39: 185–191.
- De Figueiredo E.A.P., De Sousa F.M., Guidoni A.L., Rosa P.S. (1999). Comparação de diluentes, diluições e tempo de armazenamento do sêmen sobre fertilidade, eclodibilidade e nascimento de pintos em matrizes pesadas. Rev. Bras. Zootec., 28: 1239–1244.
- D i m i t r o v S.G., A t a n a s o v V.K., S u r a i P.F., D e n e v S.A. (2007). Effect of organic selenium on turkey semen quality during liquid storage. Anim. Reprod. Sci., 100: 311–317.
- Donoghue A.M., Wishart G.J. (2000). Storage of poultry semen. Anim. Reprod. Sci., 62: 213-232.
- Douard V., Hermier D., Blesbois E. (2000). Changes in turkey semen lipids during liquid *in vitro* storage. Biol. Reprod., 63: 1450–1456.
- Douard V., Hermier D., Magistrini M., Blesbois E. (2003). Reproductive period affects lipid composition and quality of fresh and stored spermatozoa in turkeys. Theriogenology, 59: 753-764.
- Douard V., Hermier D., Magistrini M., Labbé C., Blesbois E. (2004). Impact of changes in composition of storage medium on lipid content and quality of turkey spermatozoa. Theriogenology, 61: 1–13.
- Douard V., Hermier D., Labbè C., Magistrini M., Blesbois E. (2005). Role of seminal plasma in damage to turkey spermatozoa during *in vitro* storage. Theriogenology, 63: 126–137.
- Gao D., Critser J.K. (2000). Mechanisms of cryoinjury in living cells. ILAR J., 41: 187-196.
- Garner D.L., Thomas C.A., Gravance C.G. (1999). The effect of glycerol on the viability, mitochondrial function and acrosomal integrity of bovine spermatozoa. Reprod. Dom. Anim., 34: 399–404.
- Gliozzi T.M., Zaniboni L., Cerolini S. (2011). DNA fragmentation in chicken spermatozoa during cryopreservation. Theriogenology, 75: 1613–1622.
- H a m m e r s t e d t R.H. (1995). Cryopreservation of Poultry Semen Current Status and Economics. In: Proceedings First International Symposium on the Artificial Insemination of Poultry, Bakst M.R., Wishart G.J. (eds). Poultry Sci. Assoc., Savoy, Illinois, USA, pp. 229–250.
- H a m m e r s t e d t R., G r a h a m J.K. (1992). Cryopreservation of poultry semen: the enigma of glycerol. Cryobiology, 29: 26–38.
- H an X.F., N i u Z.Y., L i u F.Z., Y ang C.S. (2005). Effect of diluents, cryoprotectants, equilibration time and thawing temperature on cryopreservation of duck semen. Int. J. Poultry Sci., 4: 197–201.

- I a f f a l d a n o N., M e l u z z i A. (2003). Effect of dialysis on quality characteristics of turkey semen during liquid storage. Theriogenology, 60: 421–427.
- Iaffaldano N., Meluzzi A., Manchisi A., Passarella S. (2005). Improvement of stored turkey semen quality as a result of He-Ne laser irradiation. Anim. Reprod. Sci., 85: 317–325.
- I a f f a l d a n o N., M a n c h i s i A., R o s a t o M.P. (2008). The preservability of turkey semen quality during liquid storage in relation to strain and age of males. Anim. Reprod. Sci., 109: 266–273.
- Iaffaldano N., Manchisi A., Gambacorta M., Di Iorio M., Rosato M.P. (2009). Effect of different sperm concentrations on the post-thaw viability and motility of turkey spermatozoa cryopreserved by the pellet method. Ital. J. Anim. Sci., 8: 760–762.
- Iaffaldano N., Rosato M.P., Paventi G., Pizzuto R., Gambacorta M., Manchisi A., Passarella S. (2010). The irradiation of rabbit sperm cells with He-Ne laser prevents their *in vitro* liquid storage dependent damage. Anim. Reprod. Sci., 119: 123–129.
- Iaffaldano N., Romagnoli L., Manchisi A., Rosato M.P. (2011). Cryopreservation of turkey semen by the pellet method: Effects of variables such as the extender, cryoprotectant concentration, cooling time and warming temperature on sperm quality determined through principal components analysis. Theriogenology, 76: 794–801.
- Iaffaldano N., Di Iorio M., Rosato M.P., ManchisiA. (2014). Cryopreservation of rabbit semen using non-permeable cryoprotectants: Effectiveness of different concentrations of low-density lipoproteins (LDL) from egg yolk versus egg yolk or sucrose. Anim. Reprod. Sci., 151: 220–228.
- Iaffaldano N., Di Iorio M., Miranda M., Zaniboni L., Manchisi A., Cerolini S. (2016). Cryopreserving turkey semen in straws and nitrogen vapor using DMSO or DMA: effects of cryoprotectant concentration, freezing rate and thawing rate on post-thaw semen quality. Br. Poultry Sci., 57: 264–270.
- K a s a i K., I z u m o A., I n a b a T., S a w a d a T. (2000). Assessment of fresh and stored duck spermatozoa quality via *in vitro* sperm-egg interaction assay. Theriogenology, 54: 283–290.
- Kotłowska M., Dietrich G., Wojtczak M., Karol H., Ciereszko A. (2007). Effects of liquid storage on amidase activity, DNA fragmentation and motility of turkey spermatozoa. Theriogenology, 67: 276–286.
- K o w a l c z y k A., Ł u k a s z e w i c z E. (2015) Simple and effective methods of freezing capercaillie (*Tetrao urogallus L.*) semen. PLoS ONE, 10: 1–11.
- K o w a l c z y k A., Ł u k a s z e w i c z E., R z o ń c a Z. (2012). Successful preservation of capercaillie (*Tetrao urogallus L.*) semen in liquid and frozen states. Theriogenology, 77: 899–907.
- Labbé C., Blesbois E., Leboeuf B., Guillouet P., Stradaioli G., Magistrini M. (2003). Technologie de la conservation du sperme chez plusieurs vertébrés domestiques: Protection des lipides membranaires, intégrité du noyau et élargissement des méthodes. Proc. Congrès du Bureau des Ressources Génétiques, La Châtre, France, 15–17.10.2002, pp. 25–33.
- L a k e P.E. (1986). The history and future of the cryopreservation of avian germplasm. Poultry Sci., 65: 1–15.
- Lemoine M., Grasseau I., Magistrini M., Blesbois E. (2011). Ability of chicken spermatozoa to undergo acrosome reaction after liquid storage or cryopreservation. Theriogenology, 75: 122–130.
- L o n g J.A. (2006). Avian semen cryopreservation: what are the biological challenges? Poultry Sci., 85: 232–236.
- L o n g J.A., B a k s t M.G. (2007). The current state of semen storage and AI technology. Proc. Midwest Poultry Federation Convention, Saint Paul, Minnesota, USA, pp. 148–152.
- Long J.A., Conn T.L. (2012). Use of phosphatidylcholine to improve the function of turkey semen stored at 4°C for 24 hours. Poultry Sci., 91: 1990–1996.
- Long J.A., Kramer M. (2003). Effect of vitamin E on lipid peroxidation and fertility after artificial insemination with liquid stored turkey semen. Poultry Sci., 82: 1802–1807.
- L o n g J.A., K u l k a r n i G. (2004). An effective method for improving the fertility of glycerol-exposed poultry semen. Poultry Sci., 83: 1594–1601.
- Long J.A., Bongalhardo D.C., Peláez J., Saxena S., Settar P., O'Sullivan N.P., Fulton J.E. (2010). Rooster semen cryopreservation: Effect of pedigree line and male age on postthaw sperm function. Poultry Sci., 89: 966–973.
- Long J.A., Purdy P.H., Zuidberg K., Hiemstra S.J., Velleman S.G., Woelders H.

(2014). Cryopreservation of turkey semen: effect of breeding line and freezing method on post-thaw sperm quality, fertilization, and hatching. Cryobiology, 68: 371–378.

- M a s s i p A., L e i b o S.P., B l e s b o i s E. (2004). Cryobiology of gametes and the breeding of domestic animals. In: Life in the frozen state, Benson E., Fuller B., Lane N. (eds). Taylor & Francis Group, London, UK, pp. 371–393.
- Mocé E., Grasseau I., Blesbois E. (2010). Cryoprotectant and freezing-process alter the ability of chicken sperm to acrosome react. Anim. Reprod. Sci., 122: 359–366.
- Neuman S.L., McDaniel C.D., Frank L., Radu J., Einstein M.E., Hester P.Y. (2002). Utilisation of a sperm quality analyser to evaluate sperm quantity and quality of turkey breeders. Br. Poult. Sci., 43: 457–464.
- Pandian C., Prabakaran R., Venukopalan K., Kalatharan J. (2011). Effect of genetic groups and cryoprotectants on preservation of turkey semen. Indian Vet. J., 88: 16–17.
- P eláez J., Bongalhardo D.C., Long J.A. (2011). Characterizing the glycocalyx of poultry spermatozoa: III Semen cryopreservation methods alter the carbohydrate component of rooster sperm membrane glycoconjugates. Poultry Sci., 90: 435–443.
- Penfold L., Harnal V., Lynch W., Bird D., Derricson S., Wildt E. (2001). Characterization of northern pintail (*Anas acuta*) ejaculate and the effect of sperm preservation on fertility. Reproduction, 121: 267–275.
- Pierson E.E., McDaniel G.R., Krista L.M. (1988). Relationship between fertility duration and *in vivo* sperm storage in broiler breeder hens. Br. Poult. Sci., 29: 199–203.
- Purdy P.H., Song Y., Silversides F.G., Blackburn H.D. (2009). Evaluation of glycerol removal techniques, cryoprotectants, and insemination methods for cryopreserving rooster sperm with implications of regeneration of breed or line or both. Poultry Sci., 88: 2184–2191.
- Rosato M.P., Centoducati G., Santacroce M.P., Iaffaldano N. (2012). Effects of lycopene on *in vitro* sperm quality and lipid peroxidation in refrigerated and cryopreserved turkey spermatozoa. Br. Poult. Sci., 53: 545–552.
- R o s a t o M.P., I a f f a l d a n o N. (2013). Cryopreservation of rabbit semen: comparing the effects of different cryoprotectants, cryoprotectants-free vitrification, and the use of albumin plus osmoprotectants on sperm survival and ferrility after standard vapor freezing and vitrification. Theriogenology, 79: 508–516.
- Saint Jalme M., Lecoq R., Seigneurin F., Blesbois E., Plouzeau E. (2003). Cryopreservation of semen from endangered pheasants: the first step towards cryobank for endangered avian species. Theriogenology, 59: 875–888.
- Santiago-Moreno J., Castaño C., Toledano-Díaz A., Coloma M.A., López-Sebastian A., Prieto M.T., Campo J.L. (2011). Semen cryopreservation for the creation of a Spanish poultry breeds cryobank: Optimization of freezing rate and equilibration time. Poultry Sci., 90: 2047–2053.
- Seigneurin F., Blesbois E. (1995). Effects of freezing rate on viability and fertility of frozenthawed fowl spermatozoa. Theriogenology, 43: 1351–1358.
- Seigneurin F., Grasseau I., Chapuis H., Blesbois E. (2013). An efficient method of guinea fowl sperm cryopreservation. Poultry Sci., 92: 2988–2996.
- S e x t o n T.J. (1974). Oxidative and glycolytic activity of chicken and turkey spermatozoa. Comp. Biochem. Physiol., 48: 39–65.
- S e x t o n T.J. (1981). Development of a commercial method for freezing turkey semen: 1. Effect of prefreeze techniques on the fertility of frozen and frozen-thawed semen. Poultry Sci., 60: 1567–1572.
- S e x t o n T.J., F e w l a s s T. (1978). A new poultry semen extender. 2. Effect of the diluent components on the fertilizing ability of chicken semen stored at 5°C. Poultry Sci., 57: 277–284.
- Siudzińska A., Łukaszewicz E. (2008). The effect of breed on freezability of semen of fancy fowl. Anim. Sci. Pap. Rep., 26: 331–340.
- Słowińska M., Liszewska E., Dietrich G.J., Ciereszko A. (2012). Characterization of proacrosin/acrosin system after liquid storage and cryopreservation of turkey semen (*Meleagris gallopavo*). Theriogenology, 78: 1065–1077.
- Słowińska M., Dietrich G.J., Liszewska E., Kozłowski K., Jankowski J., Ciereszko A. (2013). Effect of dialysis on the proacrosin/acrosin system and motility of turkey (*Meleagris gallopavo*) spermatozoa during liquid storage. Poultry Sci., 54: 661–668.

- Surai P., Wishart G.J. (1996). Poultry artificial insemination technology in the countries of the former USSR. Worlds Poultry Sci. J., 52: 27–43.
- Swain J.E., Smith G.D. (2010). Cryoprotectants. In: Fertility cryopreservation, Chian R.C., Quinn P. (eds). Cambridge University Press, New York, USA, pp. 24–38.
- Tajima A. (2013). Conservation of avian genetic diversity. Jpn. Poultry Sci., 50: 1-8.
- Tajima A., Graham E.F., Shoffner R.N., Otis J.S., Hawkins D.M. (1990). Cryopreservation of semen from unique chicken germ plasm lines. Poultry Sci., 69: 999–1002.
- Thurston R.J. (1995). Storage of poultry semen above freezing for 24–48 hours. In: First International Symposium on Artificial Insemination in Poultry, Bakst M.R., Cecil H. (eds). Poultry Sci. Assoc., Savoy, IL., pp. 107–122.
- Tselutin K., Narubina L., Mavrodina T., Tur B. (1995). Cryopreservation of poultry semen. Br. Poult. Sci., 36: 805–811.
- Tselutin K., Seigneurin F., Blesbois E. (1999). Comparison of cryoprotectants and methods of cryopreservation of fowl spermatozoa. Poultry Sci., 78: 586–590.
- Váradi E., Vegi B., Liptoi K., Barna J. (2013). Methods for cryopreservation of guinea fowl sperm. PLoS ONE, 8(4): 1–6.
- Wa t s o n P.F. (1995). Recent developments and concepts in the cryopreservation of spermatozoa and the assessment of their post-thawing function. Reprod. Fertil. Dev., 7: 871–891.
- Williamson R.G., Etches R.J., Reinhart B.S., MacPherson J.W. (1981). The effect of cooling rate before freezing and the temperature of the semen upon addition of DMSO on the fertilizing capacity of chicken semen stored at -196°C. Reprod. Nutr. Dev., 21: 1033–1042.
- Wishart G.J. (2009). Semen quality and semen storage. In: Biology of breeding poultry. Poultry Science Symposium Series, Hocking P. (ed.). London, UK, pp. 151–178.
- Wo elders H., Zuidberg C.A., Hiemstra S.J. (2006). Animal genetic resources conservation in the Netherlands and Europe: poultry perspective. Poultry Sci., 85: 216–222.
- Z a n i b o n i L., C e r o l i n i S. (2009). Liquid storage of turkey semen: changes in quality parameters, lipid composition and susceptibility to induced *in vitro* peroxidation in control, n-3 fatty acids and alpha-tocopherol rich spermatozoa. Anim. Reprod. Sci., 112: 51–65.
- Zaniboni L., Cassinelli C., Mangiagalli M.G., Gliozzi T.M., Cerolini S. (2014). Pellet cryopreservation for chicken semen: Effects of sperm working concentration, cryoprotectant concentration, and equilibration time during *in vitro* processing. Theriogenology, 82: 251–258.
- Z a v o s P.M., G r a h a m E.F. (1983). Effects of various degrees of supercoiling and nucleation temperatures on fertility of frozen turkey spermatozoa. Cryobiology, 20: 553–559.

Received: 22 XII 2015 Accepted: 5 IV 2016