The roles of reactive oxygen species and antioxidants in cryopreservation

Jia Soon Len*, Wen Shuo Darius Koh* and Shi-Xiong Tan

Diploma in Biomedical Science, School of Applied Science, Republic Polytechnic, 9 Woodlands Avenue 9, Singapore 738964

Correspondence: Shi-Xiong Tan (tan_shixiong@rp.edu.sg)

Cryopreservation has facilitated advancement of biological research by allowing the storage of cells over prolonged periods of time. While cryopreservation at extremely low temperatures would render cells metabolically inactive, cells suffer insults during the freezing and thawing process. Among such insults, the generation of supra-physiological levels of reactive oxygen species (ROS) could impair cellular functions and survival. Antioxidants are potential additives that were reported to partially or completely reverse freeze-thaw stress-associated impairments. This review aims to discuss the potential sources of cryopreservation-induced ROS and the effectiveness of antioxidant administration when used individually or in combination.

Introduction

The ability to keep an organism alive while frozen and allowing it to survive for a prolonged period of time may sound like a scene lifted directly from a science fiction movie. Although freezing complex multicellular organisms remains challenging and often faced significant obstacles during the revival of the frozen organism [1–4], reviving single-cell organisms after a prolonged period of time has been a reality for several decades. Among many cases, the ability to revive single cell prokaryotic organisms such as Escherichia coli and Treponema pallidum were demonstrated in 1913 and 1954 respectively [5,6]. Such results were also obtained from the unicellular eukaryotic organism Saccharomyces cerevisiae in 1902 [7].

In the field of research involving mammalian cells, significant progress was made when Polge et al. [8] successfully revived frozen fowl spermatozoa in 1949 and Bos taurus spermatozoa cells in 1952 using glycerol as a cryoprotective agent (CPA) [9]. The subsequent use of dimethyl sulfoxide (DMSO) as CPA, which remarkably preserved erythrocytes, was first reported in the 1950s [10] and is now a common component of cryopreservation medium. Although the ability to allow cells to be transported across the world has fostered trans-global scientific collaborations as well as independent verifications of experimental results and clinical advancements, it is frequently taken for granted. One could only imagine the hindrance to scientific advancements if the cryopreservation techniques were absent. For the past few decades, although significant advancements were made in the cryopreservation field on mammalian cells, the technique is far from perfect. Many researchers face challenges such as poor recovery [11,12], loss of functional characteristics of specific cell types [4,13,14] and, in the case of stem cell research, the inability to retain pluripotency [15,16]. In this review, we focus on the role of reactive oxygen species (ROS), a product of cellular metabolism that can be damaging to cells and how ROS contributes to the undesirable results seen after cryopreservation. We further explore current advancements in using antioxidants to negate these undesirable effects observed in cryopreservation.

Cryopreservation and ROS production

Cells have mechanisms to detoxify ROS and once these mechanisms are overwhelmed, ROS can affect various cellular functions and processes by oxidizing proteins, inducing damage to nucleic acids, and...
peroxidation of lipids [17,18]. Oxidative stress, which is the shift of redox homeostasis toward favoring formation of ROS, dictate the subsequent cellular outcomes such as cellular senescence, apoptosis and altered cellular signaling. Generally, it is known that ROS can modulate cellular survival at low concentrations and death at supraphysiological levels [19]. It is also worth noting that physiological amount of ROS can act as signaling molecules for cellular signaling events [20]. To appreciate the impact of ROS in cryopreservation, it is important to understand the different characteristics of ROS produced in cells, the intracellular sources of these ROS and how cells detoxify these damaging species. Detailed reviews on ROS can be found in published review articles [17,18,21–22] and will not be covered in detail in our current review. A short summary of the sources of ROS and enzymes involved in ROS detoxification is provided in Figure 1 and the section below. Sustained oxidative stress has been believed to be linked to senescence — a response to cellular stress [23], with many lines of evidence supporting this [23–26]. The specific effects of the individual reactive species depend on the relative levels within the cell. The effect of these species at different levels and the biological consequences are summarized in Figure 2.

ROS production has been detected in reproductive and non-reproductive cells. ROS in the form of superoxide (O$_2^•−$) which were detected in the cells of various species undergoing cryopreservation can be reduced with the addition of various antioxidants (Tables 1 and 2). O$_2^•−$ is short-lived and does not cross the mitochondrial or lipid membranes readily due to its charge [27,28]. O$_2^•−$ cannot react with most biological molecules in the aqueous environment of the cytoplasm [18]. O$_2^•−$ can be converted into hydrogen peroxide (H$_2$O$_2$) by three known superoxide dismutase (SOD) isoforms; cytosolic-localized SOD1 (Cu, Zn SOD), mitochondrial-localized SOD2 (Mn SOD) and the extracellular SOD3 (Fe SOD). The localization of the SOD isoforms are reviewed in [29]. Significantly elevated levels of O$_2^•−$ and lipid peroxidation were observed in reviving cryopreserved bull spermatozoa [30]. In alpaca sperm, higher levels of O$_2^•−$ were detected as compared with other oxidizing intermediates using fluorescent dyes dihydroethidium (DHE) for O$_2^•−$ and 2′,7′-dichlorodihydrofluorescein diacetate (H$_2$DCFDA) for ROS, which were mainly contributed by cells that are propidium iodide negative [31]. This is also seen in human retinal pigment epithelial (hRPE) cells, where apart from an increase in ROS as detected by H$_2$DCFDA only, cellular senescence as well as telomere shortening were reported to increase as a result of cryopreservation [32]. Notably, studies have shown that DMSO treatment of human embryonic stem cells (hES) increases O$_2^•−$ by two-folds while the same cells that were thawed after cryopreservation in the presence of DMSO lead to a five-fold increase in the O$_2^•−$. These data indicate that freeze-thaw stress can promote ROS generation [33]. Other known ice nucleation inhibitors such as anti-freezing protein (AFP) and polyethylene glycol (PEG) have also been known to protect against freeze thaw-induced ROS generation [34–36]. O$_2^•−$ participates in Fenton and Haber–Weiss (HW) reaction in the presence of a ferrous iron catalyst to generate hydroxyl radical (•OH) (Figure 1). •OH, unlike O$_2^•−$, can function in the aqueous state and is particularly reactive. •OH is considered the most damaging member of ROS [37] and was reported to cause oxidation of amino acids [38], and these can result in the fragmentation and disruption of protein conformation [39]. •OH can abstract hydrogen and lead to altered nucleic acid bases resulting in DNA damage [40]. Intriguingly, there are no known enzymes to detoxify •OH, despite the damage that it can cause to the cell.

With respect to H$_2$O$_2$ levels in cells during cryopreservation, total H$_2$O$_2$ levels remain largely unchanged, while mitochondrial H$_2$O$_2$ were reported to be increased in spermatozoa [30,41]. H$_2$O$_2$ has highly selective reactivity with only certain biomolecules and can cause the oxidation of thiol groups (SH). H$_2$O$_2$ is toxic at high concentrations because it can be reduced by ferrous Iron, Fe (II), into the more damaging •OH via the Fenton reaction (Figure 1) [42–44]. H$_2$O$_2$ has a long half-life which allows it to transduce signals at long ranges [37,45]. When in extracellular space, H$_2$O$_2$ can re-enter the cell through aquaporin-dependent pathways or via direct diffusion [46]. Multiple pathways such as the glutathione peroxidase-glutathione reductase and the peroxiredoxin/thioredoxin-thioredoxin reductase pathways utilize NADPH as reducing equivalent to reduce H$_2$O$_2$ to H$_2$O (Figure 1). The species of ROS and methods used for detection in different cell types used for cryopreservation are summarized in Tables 1 and 2.

**Mitochondrial ROS production**

Studies from fish [47], sheep [48] and human cells [49] have indicated that cryopreservation induced alterations and/or damages to the mitochondria. Proteins upstream in the electron transport chain (ETC) can generate ROS through the univalent donation of electrons to oxygen in the mitochondria. Sources of ROS in the mitochondria include complex I, complex II and complex III [50]. These enzymes 'leak' electrons and as a result, univariantly reduce oxygen to O$_2^•−$. Through this process, ROS in the form of O$_2^•−$, •OH and H$_2$O$_2$ are produced (Figure 1).

Factors influencing the production of ROS in the mitochondria include: tissue or cell type, oxygen tension of the extracellular environment, presence of metabolic intermediates and substrates [51], hyperoxia [50,52], the presence of a high proportion of NADH electron donors [51,53] as well as the mitochondrial membrane potential (∆ψ) and the...
Figure 1. Metabolism and sources of ROS

(A) Detoxification and metabolism of reactive oxygen/nitrogen species. (B) Sources of ROS, and localization of enzymes that counteracts ROS in the mitochondria, endoplasmic reticulum (ER), peroxisome, cytosol and the extracellular space. SOD1 is localized in both the mitochondria intermembrane space and cytosol, SOD3 is located extracellularly and SOD2 is found exclusively mostly in the mitochondria matrix. Catalase that reduces hydrogen peroxide ($H_2O_2$) into $H_2O$ is mostly located in the peroxisomes. Glutathione peroxidase (GPx) is found in the mitochondria and cytosol. Peroxiredoxins (Prx) and thioredoxins (Trx) which constitute the Peroxiredoxin–Thioredoxin (Prx/Trx) system can be found in the nucleus, mitochondria, ER, peroxisome and the extracellular environment. Electron transport chain (ETC), Cytochrome P450 family of enzymes (Cyps), xanthene oxidase (XO) and NADPH oxidases (NOX) are potential sources of $O_2^{-}$, while ERO1 and acetyl CoA oxidases (AcoX) produce $H_2O_2$. Nitric oxide synthase (NOS) is a potential source of NO. Aquaporins (Aqp) facilitate the movement of $H_2O_2$ across membranes. Single snowflake indicates ROS detected while two snowflakes indicate an implication with cryopreservation.

© 2019 The Author(s). This is an open access article published by Portland Press Limited on behalf of the Biochemical Society and distributed under the Creative Commons Attribution License 4.0 (CC BY).
| Compound | Cell type | Beneficial effects | No effect/adverse effects | Cryopreservation method |
|----------|-----------|--------------------|--------------------------|------------------------|
| 2,4-dinitrophenol (DNP) | Sperm | • Motility (↑) [76] | • Motility (N/C) [76] | 1 cm styrofoam boat on LN at 10 min [76] |
| Ascorbic acid | Sperm | • ROS• (↓) [149] | • ROS• (↓) [149] | LN vapor phase (6.5-2 cm) at 10–15 min [148] |
| | | • Viability (↑) [149] | • Viability (↑) [149] | LN vapor phase (10 cm) at 10 min [149], −20°C at 10 min + LN vapor phase at 2 h [150] |
| | | • Motility (Weak) (↑) [149] | • DNA fragmentation (N/C or ↑) [148, 150] | |
| | | • MMP (↑) [149] | • DNA damage (↓) [148, 150] | |
| | | • Apoptotic cells (↑) [149] | • Apoptotic cells (↑) [149] | |
| | | • Viability (N/C) [148] | • Viability (N/C) [148] | |
| | | • Motility (N/C) [149] | • Motility (N/C) [149] | |
| Mouse embryos | | | | |
| | | • Percentage of intact embryos, blastocyst and number of hatching blastocyst (↑) [183] | • Fetal development (N/C) [183] | Vitrification and slow freezing [183] |
| Antifreeze proteins (AFP) | Oocytes | • ROS• (↓) [35] | • ROS• (↓) [35] | Vitrification [35] |
| | | • γH2AX+ cells (↓) [35] | • γH2AX+ cells (↓) [35] | |
| | | • Viability (↑) [35] | • Viability (↑) [35] | |
| | | • Cleavage rate, blastocyst rate, blastomere count (↑) [35] | • Cells with DNA repair (N/C) [35] | |
| | | • Apoptotic blastomeres (↓) [35] | • Apoptotic blastomeres (↓) [35] | |
| | | • Improved chromosomal alignment and spindle organization [35] | • Improved chromosomal alignment and spindle organization [35] | |
| BHT | Sperm | % Motility and viability (↑) [203] | % Sperm with functional intact membrane and GPx activity (N/C) [203] | LN vapor phase (4 cm) at 15 min [203] |
| Catalase | Sperm | • ROS• (↓) [149] | • ROS• (↓) [149] | LN vapor phase (10 cm) at 10 min [149] |
| | | • Viability (↑), weak (↓) motility, MMP [149] | • Viability (↑), weak (↓) motility, MMP [149] | LN vapor phase at 20 min [125] |
| | | • (↑) Apoptotic cells [149] | • (↑) Apoptotic cells [149] | |
| | | • (↑) Apoptotic like changes, apoptotic and necrotic cells [244] | • (↑) Apoptotic like changes, apoptotic and necrotic cells [244] | |
| | | • (↓) DNA damage [149] | • (↓) DNA damage [149] | |
| | | • Motility (↑) [76] | • Motility (↑) [76] | |
| | | • % Motility and viability (↑) [244] | • % Motility and viability (↑) [244] | |
| Coenzyme Q | Oocytes | N.A [124] | Oocyte survival and fertility (N/C) [124] | Controlled rate freezing [124] |
| | | • Viability, % sperm with functional membrane and active mitochondria [154] | • Viability, % sperm with functional membrane and active mitochondria [154] | |
| | | • Weak (↑) number of abnormal cells [154] | • Weak (↑) number of abnormal cells [154] | |
| | | • Lipid peroxidation [154] [218] (PI staining was done) | • Lipid peroxidation [154] [218] (PI staining was done) | |
| | | • DNA fragmentation (↑) [154] | • DNA fragmentation (↑) [154] | |
| | | • (↑) total and progressive motility, plasma membrane integrity [218] | • (↑) total and progressive motility, plasma membrane integrity [218] | |
| Egg yolk | Sperm | N.A [116] | (↑) NO• [116] | LN vapor phase (4 cm) at 10 min [116] |
| | | | | |

Continued over
| Compound          | Cell type                  | Beneficial effects                                                                 | No effect/adverse effects                                                                 | Cryopreservation method                      |
|-------------------|----------------------------|------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|---------------------------------------------|
| Glutathione (GSH) | Sperm                      | • Fertilization rate and % cells with ability to undergo acrosome reaction (↑) [41] | • Motility-associated parameters (↑) or (N/C) [41] • DNA fragmentation (N/C) [244]       | LN vapor phase (N/I) [41] • LN vapor phase at 10 min [153] |
|                   | Germ cells enriched with   | • Lipid peroxidation, mitochondrial ROS, total ROS and intracellular ROS (↓) [41]   |                                                                                           |                                             |
|                   | spermatogonial stem cells  | • % sperm with high MMP, viability and total Progressive motility (↑) [244]         |                                                                                           |                                             |
|                   |                            | • Apoptotic like changes, apoptosis and necrosis (↓) [244]                          |                                                                                           |                                             |
|                   |                            | • Motility recovery rate (↑) [153]                                                  |                                                                                           |                                             |
|                   |                            | • Sperm DNA damage (↑) [153]                                                         |                                                                                           |                                             |
| Hemoglobin (Hb)   | Oocyte                     | • Survival and fertility (↑) [124]                                                   | N.A [124]                                                                                 | Controlled freezing [124]                   |
| Hypotaurine/Taurine| Germ cells enriched with   | • Proliferation rate and mitochondrial activity (↑) [178]                         | • Recovery of cells (N/C) [178]                                                           | Slow-freeze [178]                         |
|                   | spermatogonial stem cells  |                                                                                   |                                                                                           |                                             |
| Iodixanol         | Sperm                      | • (↑) motility (200,168), protamine, BCL2, protamine2/3 and SPACA3 expression [200] | • Acrosomal integrity (N/C) [168]                                                        | Controlled freezing [168]                   |
|                   |                            | • (↑) BAX and ROMO1 expression and cellular death [200]                             |                                                                                           | LN vapor phase (2 cm) at 15 min [200]       |
| L-carnitine       | Sperm                      | • DNA fragmentation (↑) [148]                                                       | • Motility and viability (N/C) [148]                                                      | LN vapor phase (6.5-2 cm) at 10–15 min [148] |
| L-proline         | Oocyte                     | • Survival rate (%) (↑) [217]                                                       | • Developmental parameters, apoptosis levels, spindle recovery (N/C) [217]                | Vitrification [217]                         |
| Lactoferrin/apotransferrin | Mouse embryos            | • (↑) percentage of intact embryos and blastocysts (↑) [183]                        | • Hatching blastocyst (N/C) [183]                                                        | Vitrification or slow freeze [183]          |
|                   | Sperm                      | • (↑) Fe, NO (Griess reagent system) [125]                                            | • DNA oxidation (N/C) [147]                                                               | −20°C at 8 min + LN vapor phase at 2 h [147] |
|                   |                            | • (↑) percentage of cells with functional plasma membrane [125]                     |                                                                                           |                                             |

Continued over
| Compound | Cell type | Beneficial effects | No effect/adverse effects | Cryopreservation method |
|----------|-----------|--------------------|---------------------------|-------------------------|
| Melatonin | Sperm     | (1) total antioxidant capacity, GSH concentration, functional plasma membrane cells, mitochondrial membrane integrity [199] (1) acrosomal integrity [199,240] (1) MMP [199,240] (6) SOD, catalase and GPx activity [199,240,241] (1) BCL-2, SOD2, GSTM1, NRF2, HSP90AA1, catalase and HO-1 gene expression [199] (1) Lipid peroxidation [199] [240,241] and ROS [199] [240] [241] (1) MOV7, Bax expression [240] (1) Reduced apoptotic like changes [244] | DNA fragmentation and LDH activity (N/C) [199] Total blastocyst output (N/C) [156] Viability, Bax expression and ROS (N/C) [198] | LN vapor phase (10 cm) at 1 h [199] Frozen in pellet-form on dry ice LN vapor phase at 10 min [198] (N/I) Stored in liquid nitrogen [240] Pellet freezing in LN [241] |
| Monothioglycerol (MTG) | Sperm | Mitochondrial ROS and total ROS (1) [41] Fertility and % cells with ability to undergo acrosome reaction (1) [41] Lipid peroxidation (1) [41] (1) motility recovery rate [153] (1) sperm DNA damage [153] | Motility parameters (N/C) [41] | LN vapor phase (1–5 cm) at 30 min [214] |
| NG-nitro-L-arginine methyl ester (L-NAME) | Oocyte | (1) fertility and survival (low concentration) [124] | (1) fertility and survival (high concentrations) [124] | Controlled freezing [124] |
| Quercetin | Sperm | DNA fragmentation (1) [153] Motility and recovery rate (1) [153] (1) % high MMP cells [216] | Progressive motility, acrosome and sperm plasma membrane integrity (N/C) [216] | Controlled freezing [216] LN vapor phase at 10 min [153] |
| Resveratrol | Sperm | DNA damage [150], MDA levels [219] and % high MMP cells (1) [216] SOD activity (1) [219] | Progressive motility, acrosome integrity, integrity of sperm plasma membrane (N/C) [216] | Slow cool (~20°C) at 10 min followed by LN vapor phase at 2 h [150] Controlled freezing [216] Slow cool (~20°C) at 10 min followed by LN vapor phase (N/I) at 2 h [219] |
| SOD | Oocyte | (1) fertility and survival [124] (1) motility [76] | Decrease in fertility [124] (low concentration) Motility (N/C) [76] | Controlled rate freezing [124] 1 cm styrofoam boat on LN at 10 min [76] |
| SOD | Sperm | (1) fertility and survival [124] (1) motility [76] | Total and progressive motility, DNA fragmentation, viability, % sperm with high MMP (N/C) [244] Increased late apoptotic and necrotic cells [244] | Controlled rate freezing [124] 1 cm styrofoam boat on LN at 10 min [76] |

Continued over
| Compound                  | Cell type                                      | Beneficial effects                                                                 | No effect/adverse effects                                                                | Cryopreservation method                        |
|--------------------------|-----------------------------------------------|------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|-----------------------------------------------|
| Trehalose                | Germ cells enriched with spermatogonial stem cells | • ↑ proliferation, recovery of colonies after culture and cell viability [207] • ↑ Apoptosis ([207] • ↑ Lipid peroxidation [204] • ↑ SOD and catalase activity [204] | • Formation of colonies after transplantation (N/C) [207]                                   | Slow freeze [207]                             |
| Testicular tissue        |                                               | • ↑ cell viability, GSH content and T-AOC [204] • ↑ DNA fragmentation [148] • ↑ MDA ([207] | • N.A [204]                                                                              | −20°C at 2 h, −80°C at 12 h [204]             |
| Vitamin E                | Sperm                                         | • ↑ Motility [76] • ↑ DNA fragmentation [148] • ↑ O₂•* production in live cells ([207] | • Viability and motility (N/C) [148] • Motility (N/C) [215]                             | [Controlled rate freezing] 62.3°C/min [215]    |
|                         |                                               | • ↑ SOD and catalase activity [204] • ↑ GSH content and T-AOC [204] | • -> production in live cells (N/C) [215]                                                | 1 cm styrofoam boat on LN at 10 min [76]      |
|                         |                                               |                                                                                     |                                                                                        | LN vapor phase (6.5-2 cm) at 10-15 min [148]  |
| Zinc oxide nanoparticles | Sperm                                         | • DNA damage and lipid peroxidation [155] • ↑ Mitochondria integrity, % sperm with ability to undergo acrosome reaction and capacitation [220] • ↑ Motility ([220] | • Sperm motility and ability to undergo the acrosome reaction (N/C) [155] | (N/I) Stored at −196°C [155]                  |
| Zinc sulfate             | Sperm                                         | • DNA damage ([220] • ↑ Mitochondria integrity, % sperm with ability to undergo acrosome reaction and capacitation [220] • ↑ Motility ([220] | • Motility (N/C) [220]                                                              | LN vapor phase at 5 min [220]                 |
| Trolox (Vitamin E analog)| Ovarian tissue                                 | Viable follicles ([1] [81] • BMP4, BMP15, CTGF, GDF9, KI expression ([1] [81] | HSP70, ERp60, SOD1 and ERP29, AMH expression (N/C) [81]                              | 2°C/min from 20 to −7°C; cooled at 0.3°C/min to −30°C, into LN (−196°C) [81] |

**Table 1 Antioxidants and their effects on cryopreserved reproductive-associated cells/tissues (Continued)**

Abbreviations: BHT, butylated hydroxytoluene; LN, liquid nitrogen; MDA, malondialdehyde; MMP, mitochondrial membrane potential; N.A, not-applicable; N/C, no changes/no effect. "-, no effects have been reported. ([1] and ([2]), represent a significant increase or decrease respectively.

^, denotes cases where effects are context dependent and due to factors such as cell quality and species.

Method employed for detection of ROS and Oxidative biomarkers are denoted by alphabetical superscripts 'a' to 'u':

- a, H₂DCFDA.
- b, H₂DCFDA/Propidium Iodide (Pi).
- c, 4,5-diaminofluorescein diacetate (DAF-2DA).
- d, DAF-2DA/Ethidium Homodimer -.
- e, 8-OH-G.
- f, BODIPY 581/591 C11.
- g, BODIPY 581/591 C11/Propidium Iodide (Pi)-
- h, Bromopyrogallol Red.
- i, DHE/Sytox-.
- j, DHE.
- k, Dihydrorhodamine(DHR) 123/Propidium Iodide (Pi)-
- l, DHR 123.
- m, 2,4-dinitrophenylhydrazine (DNPH) assay.
- n, Fox2-modified method.
- o, Formamidopyrimidine-DNA glycosylase-sensitive comet assay.
- p, Griess reagent method.
- q, MitoPY1/SYTOX--.
- r, PF6-AM/SYTOX.
- s, Peroxy Green 1 (PG1).
- t, Thiobarbituric acid reactive substances (TBARS) assay.
- u, commercial or obscure ROS detection techniques.
### Table 2 Antioxidants and their effects on non-reproductive cell types/tissues

| Compound | Cell type | Beneficial effects | No effect/adverse effects | Cryopreservation method |
|----------|-----------|--------------------|--------------------------|------------------------|
| Ascorbic acid | Bone-marrow mononuclear cells | • Clonogenic parameters (↑) [221] (murine model) | • Viability and clonogenic parameters (human model) (N/C) [221] | Controlled rate freezing [221] |
| Astragalosides | Pancreatic islets | • Restored blood glucose to normal [232]  
• Insulin expression after transplantation (↑) [232] | N.A [232] | Slow-freeze [232] |
| BHT | Blood cells | • Loss of HUFA (↓) [184] | N.A [184] | Chromatography paper at −20°C [184] |
| BHT + ascorbic acid | Hepatocytes | • Post-thaw albumin production (↑) [235]  
• Urea synthesis, ammonia clearance and cell proliferation (N/C) [235]  
• Apoptosis associated DNA fragmentation (N/C) [235] | Induced LDH release (↑) [235]  
• Lipid peroxidation (N/C) (when used with glycerol as a CPA) [158] | Slow freeze [235] |
| Catalase | Mononuclear cells | • Clonogenic parameters (↑) (murine model) [221] | Viability, clonogenic parameters (human model) (N/C) [221] | Controlled rate freezing [221] |
| Catalase + Trehalose | Hematopoietic cells | • DCF fluorescence intensity (↑) [233]  
• Number of DCF+ cells (↑) [233]  
• (↑) CFU [233,234]  
• (↑) viability [233]  
• Apoptosis (↓) [233]  
• (↑) responsiveness to migratory homing associated cytokines, expression of homing-associated receptor and adhesion capacity [234] | N.A [233,234] | Controlled rate freezing [233,234] |
| Consumption of blueberries by PBMC donors | Peripheral blood mononuclear cell | • DNA oxidation (↓) [144] | DNA damage induced by H₂O₂ in cryopreserved cells (N/C) [144] | Slow freeze [144] |
| Deteroxamine | Blood cells | • Loss of HUFA (↓) [184] | N.A [184] | Chromatography paper at −20°C [184] |
| Glutathione (GSH) | Embryonic stem cells | • ROS(•) (↑) [247]  
• Viability (↑) [247] | N.A [247] | (N/I) Stored in −80°C freezer at 24 h [247] |
| Embryogenic callus | • Post-thaw survival, GSH, ascorbic acid levels, SOD and peroxidase activity (↑) [174]  
• *OH, H₂O₂, O₂−, and MDA levels (↑) [174]  
• E-Cadherin cell adhesion proteins (N/C) [236] | At high concentrations, survival (↑) or (N/C) [174]  
• Catalase activity (N/C) [174] | Slow freeze [236] |
| Pancreatic islets | • MDA (↑) [222]  
• Islet morphometry and glucose clearance rate (↑) [222] | Islet insulin secretion (N/C) [222] | Slow freeze [222] |
| Peroxiredoxin | Murine hepatocytes | • Viability (↑) [236]  
• Integrin-β1 and β-catenin cell adhesion proteins (↑) [236]  
• Urea secretion (↑) [236]  
• NOx (•) (↑) [236]  
• ROS(•−) (↑) [236]  
• O₂− (•) (↑) [236] | E-Cadherin cell adhesion proteins (N/C) [236] | Slow freeze [236] |
| Murine insulinoma | • Viability (↑) [236]  
• Insulin secretion (↑) [236]  
• NOx (•) (↑) [236]  
• O₂− (•) (↑) [236] | ROS(•) (N/C) [236] | Slow freeze [236] |
| Polyethylene glycol (PEG) | Human embryonic stem cells | • ROS(•) (↑) [36]  
• Alleviation of F-actin levels [36]  
• Cell viability (N/C) [36] | Slow freeze [36] |
| S-Adenosylmethionine | Hepatocytes | • GSH content and cellular viability (↑) [177] | N.A [177] | Slow freeze [177] |
| Salicic acid | Red blood cell | • (↑) protein carboxylation [158]  
• Lipid peroxidation (N/C) (when trehalose was used as a CPA) [158] | Lipid peroxidation (N/C) (when used with glycerol as a CPA) [158] | N.I [158] |
| SOD | Bone-marrow mononuclear cells | • N.A [221] | Post-thaw recovery (N/C) [221] | Controlled rate freezing [221] |

Continued over
| Compound               | Cell type     | Beneficial effects                                                                 | No effect/adverse effects   | Cryopreservation method |
|------------------------|---------------|-------------------------------------------------------------------------------------|-----------------------------|-------------------------|
| Trehalose              | Dendritic cells | • Preserved cell function and phenotype [205]                                       | • N.A [205]                 | Controlled rate freezing [205] |
|                        |               | • ↑ viability [205]                                                                |                             |                         |
|                        |               | • Maintained MMP and cytoskeleton integrity [205]                                  |                             |                         |
|                        |               | • ↓ apoptosis, BIM-1 and CASP9 expression [205]                                    |                             |                         |
|                        |               |                                                                              |                             |                         |
|                        | Hepatocytes   | • ↑ albumin secretion, plating efficiency and viability [206]                     | • EROD and ECOD activity, proliferation, LDH, urea levels (N/C) [206] | Controlled rate freezing [206] |
|                        |              | • ↓ AST activity [206]                                                            |                             |                         |
|                        | BM-MNC        | • (↑) Clonogenic parameters (murine and human models [221])                      | • N.A [221]                 | Slow controlled rate freezing [221] |
|                        |              |                                                                              |                             |                         |
| Wheat proteins or     | Hepatocytes   | • ↑ attachment efficiency and viability [227]                                     | • N.A [227]                 | Slow freeze [227]       |
| Lipocalins             |              | • Restoration of cytochrome P450 isoform activity to fresh cells levels [227]      |                             |                         |

Abbreviations: AST, aspartate aminotransferase; CFU, colony forming units; DCF , 2',7'-dichlorofluorescein; BHT, butylated hydroxytoluene; HUFA, highly unsaturated fatty acid; LN, liquid nitrogen; MDA, malondialdehyde; N.A, not-applicable; N/C, no changes/no effect. -, no effects have been reported. (↑) and (↓), represent a significant increase or decrease respectively. ∧, denotes cases where effects are context dependent and due to factors such as cell quality and species.

Method employed for detection of ROS and Oxidative biomarkers are denoted by alphabetical superscripts 'a' to 'u':

a, H2DCFDA.
b, H2DCFDA/Propidium Iodide (Pi).
c, 4,5-diaminofluorescein diacetate (DAF-2DA).
d, DAF-2DA/Ethidium Homodimer-

e, 8-OHG.
f, BODIPY 581/591 C11.
g, BODIPY 581/591 C11/Propidium Iodide (Pi)-
h, Bromopyrogallol Red.
i, DHE/Sytox-
j, DHE.
k, Dihydrorhodamine (DHR) 123/Propidium Iodide (Pi)-
l, DHR 123
m, 2,4-dinitrophenylhydrazine (DNPH) assay.
n, Fox2 modified method.
o, Formamidopyrimidine-DNA glycosylase-sensitive comet assay.
p, Griess reagent system.
q, MitoPY1/SYTOX-
r, PF6-AM/SYTOX-
s, Peroxy Green 1 (PG1).
t, Thiobarbituric acid reactive substances (TBARS) assay.
u, Commercial or obscure ROS detection techniques.

pH gradient [54–56], which are constituents of proton-motive force, Δp. The multitude of mitochondria inducers underlie the fact that multiple mechanisms can affect the genesis of mitochondrial ROS in the ETC (reviewed in [50,51,56]).

The MMP or Δψ is a parameter widely used to assess mitochondrial function. Δψ was reported to be altered in thawed cells following cryopreservation [57–59]. Reduction in Δψ in certain cases, such as a mild decrease, is associated with a decline in ROS levels while an increase in Δψ has been noted to promote ROS formation in rat mitochondria isolated from brain [60] and heart muscles [61]. These studies indicated that maintenance of the Δψ is an important aspect to prevent ROS-induced oxidative stress during cryopreservation.

Hyper-polarization of the Δψ can favor ROS generation [62], which is believed to be a result of a reduction in electron transfer [63]. Depolarization of Δψ can be induced by ROS, which impairs oxidative phosphorylation and amplifies ROS generation [64]. Loss of Δψ was reported in cryopreserved human oocytes [57], buffalo sperm [65,58], nucleus pulposus-derived mesenchymal stem cells [66], murine embryos [67], Meleagris gallopavo spermatozoa [68], koala spermatozoa [69] and porcine hepatocytes [59], although a transient elevation in Δψ was reported in murine oocytes after freeze-thawing [70]. Opening of mitochondrial permeability transition pore (mPTP), which involves the formation of a 'hole' in the inner mitochondrial membrane (IMM) is known to lead to the dissipation of
Figure 2. Effects of different levels of reactive oxygen/nitrogen species on cellular biomolecules

Protein can react with ONOO$^-$, H$_2$O$_2$, NO$^*$, *OH and aldehydes such as 4-Hydroxynonenal (4-HNE) can react with protein side chains (e.g. amino acids such as lysine). The formation of oxo-histidine and disulfide bonds are mostly reversible and mediate redox signaling under mild oxidative stress and may not be deleterious. High level of ROS lead to protein aggregation, denaturation and fragmentation. Mitochondrial/nuclear DNA can react with O$_2$•$^-$, ONOO$^-$ and •OH. Mutations and double/single-strand breaks mediated by ROS are minimized by the DNA-Damage Response (DDR). Proteins such as p53, RAD51 and yH2AX are DDR constituents involved in cryopreservation. Severe oxidative stress can overwhelm the DDR, resulting in mutations and double/single strand breaks. Lipids can react with ONOO$^-$ and *OH to cause lipid peroxidation and form lipid peroxides (LPOs). LPOs can decompose into aldehydes (Ald) such as 4-HNE and malondialdehyde (MDA). At low levels of ROS, cells are quiescent. Moderate levels of ROS facilitates beneficial redox signaling to modulate cellular survival, growth and division. Overwhelming levels of ROS can initiate cell death.

the Δψ as well as an elevation in ROS levels [64]. Opening of mPTP leads to dissipation of the Δψ, mitochondrial swelling, ATP depletion, relocalization of pro-apoptotic molecules and elevated ROS levels [71,72].

The involvement of mPTP in cryopreservation has been implicated in the study showing that inhibition of mPTP by bongkrekic acid successfully reduced cryopreservation-induced apoptosis in stallion spermatozoa [73]. mPTP opening has been known to enhance H$_2$O$_2$ production through conformational alterations to complex I of ETC [74], depletion of ROS-scavengers as well as intensifying production of ROS from Krebs cycle oxidoreductases [75]. Opening of mPTP is known to be induced during oxidative stress and ROS-mediated alterations to mPTP regulators and components were suggested to be responsible for this. Indeed, it was found that the mild uncoupling agent 2,4-dinitrophenol, which normally reduces ROS, improved motility in sperm with low cryopreservability [76] while
the antioxidant, monothioglycerol was found to reduce mitochondria ROS as well as increase fertility and the percentage of cells with the ability to undergo acrosome reaction [41]. ROS-mediated mitochondrial permeabilization involved oxidative attack on the protein thiol groups on the mitochondrial membrane. This may give rise to protein aggregates due to thiol groups cross-linking after being oxidized [77,78].

**Protein folding in the endoplasmic reticulum and ROS production**

Cryopreservation of cells was known to perturb the homeostasis of the endoplasmic reticulum (ER) [79–82] and ER is a known source of ROS [83,84]. The ER facilitates the proper folding and addition of some post-translational modifications to proteins in the secretory pathway. Accumulation of misfolded proteins in the ER could occur under conditions that perturb ER homeostasis, also known as ER stress. Increased protein synthesis is one such condition. The unfolded protein (UPR) response promotes an adaptive response against ER stress by increasing machineries for protein degradation and protein folding as part of an effort to restore ER homeostasis. The molecular mechanism on how UPR are activated has been comprehensively reviewed by [85,86]. UPR is activated via one of the three membrane-bound transducing receptors (ATF6, PERK, IRE1\(\alpha\)), these three sensors thus constitute the three branches of the UPR signaling pathways [85,86]. It was observed that SOD1 and the ER stress marker ERP29 gene expression were significantly up-regulated in response to freeze-thaw stress in primate ovarian tissue [81]. In yeast, genes expression for protein chaperones such as SSA4, HSP26, HSP42 were found to be up-regulated in response to freeze-thaw stress when cells were frozen in the absence of cryptoprotectants [87]. In mammals, all three arms of the UPR may be activated during cryopreservation. The XBP-1 protein levels were elevated in vitrified mouse oocytes [79] and maturing oocytes exposed to delipidated serum were more susceptible to cryopreservation-induced ER stress [80]. Intriguingly, the handling of oocytes, itself, was sufficient to activate the IRE1\(\alpha\) arm [88], highlighting the vulnerability of oocytes to cope with stress during the cryopreservation process.

In addition to its homeostatic role, sustained induction of the UPR in response to severe ER stress caused by a multitude of factors can antagonize cellular survival, resulting in cell death [89]. The activation of the UPR has been implicated in the production of at least two species of ROS: \(O_2^{•−}\) and \(H_2O_2\) [90–92] and these ROS were postulated to be an event preceding cell death. The source of \(H_2O_2\) may be attributed to oxidative folding via the protein disulfide isomerase (PDI)-ER oxidoreductase (ERO) relay or the cytochrome P450 (CYP) family of enzymes. The PDI-ERO1 pathway has been demonstrated to produce ROS in the form of \(H_2O_2\) [93–95]. Activation of the PERK-arm of the UPR could lead to downstream ERO1\(\alpha\) activation, \(H_2O_2\) production and mediates the feeding of calcium into the mitochondria which could promote \(O_2^{•−}\) production and apoptosis [92]. Interestingly, a yeast strain deleted for genes encoding for catalases and glutathione were hypersensitive to exogenous \(H_2O_2\), but are not sensitive to \(ERO1\) overexpression [96], indicating that there could be other, more potent sources of \(H_2O_2\) in the ER, or that cytosolic or mitochondria pool of \(H_2O_2\) are isolated from the ER. Studies have also indicated an ERO1-independent source of \(H_2O_2\) in the ER [97,98], suggesting the PDI-ERO1 pathway may not be the sole source of \(H_2O_2\) in the ER. Other possible pathway that could be activated through sustained UPR activation that may lead to mitochondrial ROS generation is through the dimerization of IRE1\(\alpha\), which activates the JNK-SAB axis to initiate cellular death [99].

**Nitric oxide synthase and NADPH oxidase**

Although not considered ROS, nitric oxide (NO\(•\)) and peroxynitrite (ONO\(^{-}\)) are free radicals. NO\(•\) can diffuse through the cell membrane. In vivo, NO\(•\) is not highly reactive to most biomolecules. However, NO\(•\) can react with metal complexes to form metal nitrosyls. NO\(•\) react with \(O_2^{•−}\) to form more damaging species, such as ONOO\(^{−}\) which is thought to occur mostly in the hypoxic regions of the cell [100]. ONOO\(^{−}\) can be detoxified by enzymes such as peroxiredoxins and glutathione peroxidase [101] (Figure 1). Unlike NO\(•\), ONOO\(^{−}\) are strong oxidants capable of causing oxidative damage, nitration and S-nitrosylation of proteins [102,103]. In vivo, nitric oxide (NO\(•\)) is produced by the family of nitric oxide synthase (NOS) which consist of three isoforms: neuronal NOS or NOS1 (‘neuronal’ NOS/nNOS), NOS2 (‘inducible’ NOS/iNOS) and NOS3 (‘endothelial’ NOS/eNOS). NOS typically catalyzes the formation of NO\(•\) and citrulline from arginine and oxygen. Most NOS isoforms are usually regulated by calmodulin and calcium, and require the cofactors NADPH, FAD, Flavin mononucleotide (FMN) and tetrahydrobiopterin (BH4) [104,105]. Similar to ROS, NO\(•\) regulates cell death and survival [106–110]. NO\(•\) is essential for proper cellular physiological function such as vasodilation [111] as well as regulating immunosuppression [112] and tissue repair in mesenchymal stem cells (MSCs) [113]. Conversely, NO\(•\), can interfere with hemopoiesis [114]. Moderate levels of NO\(•\) can initiate capacitation [115] and is essential for motile functions in sperm [116,117].
During cryopreservation, NOS activation or NO\(^*\) production was observed in cryopreserved heart valves [118] and sperm [30,116,119]. While NO\(^*\) itself has not been found to be significantly increased by freeze-thaw stress in RBC, the product of nitric oxide nitrosylation, S-nitrosohemoglobin was found to be increased by freeze-thaw stress [120]. At high levels of NO\(^*\), sperm functions can be antagonized [116,121–123]. When cryopreserving sperms, the use of low concentrations of NOS inhibitor, NG-nitro-l-arginine methyl ester (l-NAME) [124], and anti-nitrosative agents such as hemoglobin [124] and lactoferrin [125] has been found to improve membrane functionality, survival and/or fertilization, indicating that reducing NO\(^*\) may improve assisted reproductive technology outcomes especially since NO\(^*\) was elevated in post-thawed cells. It should however be noted that the use of high concentrations of l-NAME was found to impair sperm function [124].

NADPH oxidases (NOXs) are a family of seven-membered enzymes that are highly regarded due to their role as a major non-mitochondrial ROS generator. NOX enzymes generate ROS, primarily O\(_2\)\(^{−}\), by catalyzing the transfer of one electron across the membrane from the electron-donating NADPH to the electron acceptor oxygen, thus reducing oxygen to form O\(_2\)\(^{−}\). Exceptions are NOX4, DUOX1 and DUOX2 of the NOX/DUOX family, which was documented to produce mainly H\(_2\)O\(_2\) [126]. While all seven members of the NOX family are found to be located to the plasma membrane, specific NOX isoforms such as NOX4, NOX5 and DUOX2 have also been detected at ER, with NOX4 residing at other subcellular locations including the mitochondria and the nuclear membrane [126,127]. Apart from the mitochondria and ER, the peroxisome is another source of intracellular ROS, which harbors pro-oxidant enzymes such as acyl-CoA oxidase (ACOx) and xanthine oxidases (XOs) [128].

Our current understanding of the activation of NOX includes a collection of inducers which can be sorted into three main categories namely: chemical, biological and physical [129]. With respect to cryopreservation, NOX activation induced by chemical and physical inducers are particularly relevant and worth noting. Physical inducers are a broad collection of inducers including temperature [130], osmotic stress [131] and pH changes [132] which are documented NOX-inducers, that are coincidentally generated during cryopreservation [133]. During cryopreservation, the addition and removal of cryoprotectants, as well as freeze-thawing have been proven to subject cells to osmotic stress [134]. Extracellular ice formed during cryopreservation puts the cell through hypertonic conditions as the solute concentration elevates in the unfrozen extracellular portions. As a result, cells shrink as water leaves the cell to re-establish the equilibrium of solute concentration across the cell. The reverse is also true for thawing during cryopreservation, in which this time, cells are put through hypotonic conditions which lead to movement of water into the cell, consequently, cell swelling. Swelling of cells under hypotonic condition, however, was viewed as more pernicious due to the elevation in ROS levels following cryopreservation in the case of stallion sperm [135].

In astrocytes, hypo-osmotic swelling leads to an increase in ROS as well as phosphorylation of p47\(^{phox}\) and that, apocynin, an NOX inhibitor abrogated such effects [136]. Moreover, cortical brain slices of mice with p47\(^{phox}\) knockout failed to show elevated ROS levels as observed in wild-type mice suggesting hypo-osmotic swelling results in p47\(^{phox}\)-NOX-dependent generation of ROS [136]. In agreement with this, supporting evidence from skeletal muscles, in which osmotic stress leads to localized increase in Ca\(^{2+}\) in the cytosol, termed as ‘calcium spark’ and an elevation in ROS levels have further substantiated this viewpoint. In addition to this, treatment of skeletal muscle cells with NOX inhibitors, apocynin and diphenyleneiodonium, reversed this effect. The exclusion of extracellular Ca\(^{2+}\) restrained the increase in levels of ROS as well as calcium spark and the inhibition of Ca\(^{2+}\) release from the sarcoplasmic reticulum by the inhibitors, ryanodine and thapsigargin were able to further reduce ROS levels [131].

Taken together, the above observations suggested that NOX activation via osmotic stress may be dependent on Ca\(^{2+}\) release from the sarcoplasmic reticulum. The Ca\(^{2+}\) could then influx into mitochondria from osmotic stress, leading to induction of NOX activity [131]. Thus, it could be inferred from the above studies that cryopreservation may induce NOX activation. Whether NOX inhibitors can abrogate the ROS generated in post-thawed cells remains to be investigated. Current findings indicate that NOX activation during cryopreservation could be a potential target to reduce ROS-induced damage in cells.

**DNA damage, protein oxidation and lipid peroxidation in cryopreservation**

Detection of oxidative damage in cryopreserved cells is a valuable measurement to determine the degree of damage. Many consequences of ROS-induced damages can be credited to lipid peroxidation [137], DNA damage [138] and protein oxidation [139,140] (Figure 2). Methods used for measurement of these damages are reviewed by [141].

Cryopreservation significantly increased DNA damage in cells as assessed by the comet assay or DNA fragmentation. Activation of DNA damage repair (DDR) constituents such as p53 [33,142], γH2AX and RAD51 [143] were observed during slow-freeze and/or vitrification. DNA oxidation was increased in cryopreserved human peripheral
blood mononuclear cells (PBMCs), indicating oxidative damage has occurred in these cells [144]. In contrast, PBMCs from donors who consumed wild blueberries rich in antioxidants has been reported to have significantly lower DNA oxidation following cryopreservation. Whereas antioxidants may reduce DNA oxidation during freeze-thawing, l-carnitine, an antioxidant [145,146], has however, failed to reduce DNA oxidation in thawed human spermatozoa in vitro [147]. Addition of compounds with known antioxidant properties such as vitamin C [148–150], vitamin E [148,151], resveratrol, [150], β-mercaptoethanol [152], taurine, hypotaurine [148], glutathione (GSH) [153], coenzyme Q [154], quercetin [153], zinc oxide nanoparticles [155], catalase [149] monothioglycerol, glutathione [153] and melatonin [156] have been reported to significantly reduce DNA damage in cryopreserved cells (Table 1).

Protein oxidation, as determined by protein carbonylation were detected in cryopreserved cells. Freezing stress was characterized to lead to the formation of carbonyl groups in intact and homogenized tissue [157]. RBCs cryopreserved with glycerol or trehalose were found to have increased ROS accumulation and protein oxidation. Supplementation of the antioxidant Salidroside ameliorated this effect [158]. Protein oxidation and increased ROS was also detected in cryopreserved sperm cells [159]. Lipid peroxidation can be due to the effect of *OH and ONOO− [160,161]. Increase in lipid peroxidation was observed in tissue specimens stored at −20°C [162]. Lipid peroxidation as measured by either malondialdehyde (MDA) or 4-Hydroxynonenal (4-HNE) were detected in cryopreserved red blood cells [158], sperm [163–166] and hepatocytes [167]. Notably, the product of lipid peroxidation 4-HNE is extremely reactive, which allows it to react with DNA and proteins. Antioxidants such as iodixanol can reduce lipid peroxidation in cryopreserved buffalo semen [168].

Effectiveness of antioxidants in preventing cryoinjury: lesson learnt so far

Endogenous defense mechanisms and effects of inhibitors on ROS-generating sources in cryopreservation

Transcriptomic studies have shown that many antioxidant genes such as SOD1, cytosolic catalase T (CTTI) and glutaredoxin-1 (GRX1) were induced in the model eukaryote Saccharomyces cerevisiae, also commonly known as the baker’s yeast or brewer’s yeast [169]. These studies indicated the importance of the role of antioxidants in mitigating freeze-thaw stress after cryopreservation [169]. Intriguingly, genetic screening of yeast mutants defective for different antioxidant genes highlighted that not all antioxidants contribute equally in their ability to protect cells from freeze-thaw stress [87,170]. It was found that yeast strains deleted for SOD1 and SOD2 were particularly sensitive to freeze-thaw stress, while single deletion of catalase and glutathione peroxidase were not as sensitive [170]. In mammals, both vitrification and slow freezing were found to up-regulate SOD gene expression and increase proteins levels in murine oocytes [171], embryos [172] and testicular tissue [142]. Furthermore, the addition of O2•− scavenging agent MnCl2 rescued cells deleted for the SOD1 gene [170]. Collectively, these studies highlight the importance of the SOD genes in cryopreservation of various cell types.

Besides SOD, the reduced GSH regeneration system or the pentose-phosphate shunt for NADPH production were up-regulated in S. cerevisiae during freeze-thaw [169]. Given that GSH is the most abundant antioxidant in almost all cell types [173], it is therefore not surprising that the glutathione cycle is required for freeze-thaw tolerance. Studies where spermatozoa were administered with GSH or thiols were demonstrated to modestly reduce ROS [174], increase the motility of spermatozoa [41,175] and the developmental competence of mouse oocytes [176]. In addition, the GSH and cysteine precursor, S-adenosylmethionine, increased the total GSH levels and the viability of cryopreserved cells. While the supplementation of S-adenosylmethionine lead to significantly lower MDA levels in cold-stored rat hepatocytes, it was however not determined in the cryopreserved cell group [177]. The proliferation of spermatogonial stem cells was however noted to be unaffected by administration of glutathione [178].

Interestingly, one of the more oxidizing environment in the cell is the ER, where the reduced GSH to oxidized glutathione (GSSG) ratio is 3:1 as compared with the cytosol where the ratio is 100:1 [179]. Perturbation of ER homeostasis was known to trigger the UPR. The induction of UPR coincides with a reduction in the developmental competence and modest reductions in survival of cryopreserved cells, which can be improved by supplementation of ER stress inhibitor TUDCA [79,80,82]. The use of Trolox, a water-soluble analog of vitamin E, increased antioxidant capacity, prevented ER stress and improved the viability of ovarian tissues. This indicates a role for both oxidative stress and ER stress during cryopreservation [81]. Intriguingly, it was found that an inhibitor that prevents ER stress-induced apoptosis, Salubrinal, did not improve development and viability of bovine blastocyst [180], indicating that preventing ER stress-induced cell death alone may not be sufficient to prevent cryopreservation-induced damage. However, it is
worthy to note that in this specific case, the viability of the blastocyst is close to 100% in both control and Salubrin treated groups [180].

Gene expression encoding for proteins which regulate or sequester the availability of Fenton reaction initiators are up-regulated in transcriptomic studies in freeze-tolerant animals or in yeast undergoing freezing stress [181,182]. As antioxidants, iron chelators such as deferoxamine, lactoferrin and transferrin were found to limit NO\(^{−}\) production and improve cellular parameters affected by cryopreservation-induced oxidative stress [125,183,184]. Deferoxamine was found to prevent loss of highly unsaturated fatty acids in RBCs stored at −20°C for a shorter period time as compared with the lipophilic free radical scavenger butylated hydroxytoluene (BHT) [184]. Interestingly, supplementation of transferrin, ascorbic acid and a combination of both compounds generally improved the percentage of intact embryos. However only when ascorbic acid was used alone did the number of hatching blastocysts appreciably increase [183]. These studies revealed the complexity of the outcome of cryopreserved cells when using antioxidants as a supplement for cryopreservation.

Apoptosis as an adversity after cryopreservation
The efficiency of the cryopreservation process is still partially compromised due to several factors. Reduced cell viability, increased senescence and impaired cellular functions are among the most widely reported adversities associated with oxidative stress generated during cryopreservation of cells. In spermatozoa, freeze-thawing during cryopreservation greatly reduced cell viability accompanied by a range of structural abnormalities and damages, presumed or found to be a consequence of oxidative stress [185–187]. Supplementation of antioxidants into the cryopreservation media generally yielded good cell viability, indicating that oxidative stress plays a role in inducing cellular death during cryopreservation [147,188,189]. Cryopreservation led to re-localization of phosphatidylserine from the inner to the outer leaflet of plasma membrane, a signal displayed by cells undergoing cell death [166,190]. Caspase activation, which is well-known to be involved in mediating the apoptotic cascade, were also observed in cryopreserved sperm cells [191–193]. The use of caspase inhibitors significantly improved the viability of hepatocytes and human embryonic stem cells after cryopreservation [194–196], indicating that preventing caspase activation can be a plausible approach to improve cell viability.

Some antioxidants may exert their effects through modulation of genes responsible for pro-survival, apoptosis and/or oxidative stress. Melatonin, iodixanol, catalase and vitamin E can up-regulate anti-apoptotic genes such as Bcl2L1 (Bcl-xL) and Bcl-2, while down-regulating pro-apoptotic genes such as BAX/Bax [197–200]. With regard to melatonin, Deng et al. [199] and Chen et al. [198] observed different outcomes on BAX/Bax modulation. This difference could be attributed to the cell type used in each study. Apart from modulating genes responsible for cell survival, the pro-oxidative genes, ROMO1 (in canine) and NOX5 (in humans) have also been reported to be down-regulated by the administration of iodixanol [200] and melatonin, respectively [199]. The use of melatonin in cryopreservation has been noted to increase human antioxidant genes, such as NRF2 and SOD2 among others as shown in Table 1. It remains unclear if the up-regulation of antioxidant genes after antioxidant treatment provides direct benefit, if any, to protect cells against cryopreservation-induced ROS damage. Trehalose [201,202] and BHT [203] were reported to reduce lipid peroxidation [204] in testicular tissue and spermatozoa, respectively, generally enhance total antioxidant capacity, improve cellular viability [204–207] and reduce apoptosis [205,207]. Decrease in mitochondrial membrane potential (ΔΨ) has been observed in cells stimulated by apoptotic stimulus [208–211], which is also seen in thawed cells after cryopreservation [65]. The reduction in ΔΨ is, however, prevented through antioxidant administration [212–216]. Administration of amino acid with antioxidant properties such as l-proline is one such case where it reduces ROS levels as well as increases ΔΨ [217]. These studies indicate that supplementing antioxidants and/or factors that modulate the process of cell death can be a potential solution to reduce cryopreservation-induced cell death.

Context-dependent effects of antioxidants in cryopreservation
The effectiveness of antioxidants in ameliorating functional parameters during cryopreservation is also dependent on the cell type used as well as the integrity of the cells prior to cryopreservation. This could be observed in the cryopreservation of sperm cells from different organisms. In one example, Dong et al. reported that the beneficial effects of SOD administration were only seen in sperm with low post-thaw survivability [76]. This is also observed for other antioxidants namely coenzyme Q [218], resveratrol [219], zinc sulfate [220], ascorbic acid [150], catalase [76] and 2,4-dinitrophenol [76]. These studies concluded that the effectiveness of antioxidants was dependent on sperm quality. While vitamins C and E may generally reduce DNA damage of spermatozoa from human and Gilt-head seabream (Sparus aurata) [148,149,151], these antioxidants can increase DNA damage in cryopreserved sperm from European seabass (Dicentrarchus labrax) [148] suggesting the possibility that antioxidants ameliorate freeze-thaw stress in a
species-dependent manner. In cases where trehalose was used to cryopreserve spermatogonial stem cells, while the proliferation capability of such cells was increased in vitro, this did not translate to a real improvement in the number of colonies formed when such cells were subsequently transplanted [207]. Such disagreement between in vitro and in vivo results can also be seen when mouse embryos were incubated with ascorbic acid prior and post-cryopreservation, where the number of normal fetuses was unchanged despite notable improvements such as embryo intactness as well as blastocyst stage [183]. Consistently, there have also been reports in the literature where cellular functionality saw no improvement after antioxidant supplementation for cryopreservation [41,178,219,221,222]. Table 1 is a summary of the effect of antioxidants on cryopreserving reproductive cells and embryos.

**Potential practical applications of antioxidants and their effects on cellular function**

Hemopoietic stem cells, hepatocytes and islet cells, all possess enormous potential when transplanted. In some studies, it has been reported that the ability to synthesize proteins [223] such as insulin [224,225] or albumin [226], metabolize xenobiotics [226,227], transplantation potential [226–228] and clonogenic potential [229–231] may either be lost or impaired via the process of cryopreservation. Such impairments or undesirable outcomes of cellular functionality have been partly improved through administration of antioxidants. These include ascorbic acid [221], astrogalosides [232], taurine [222], hypotaurine [178], vitamin E [76], catalase [221], trehalose [205–207], combination of catalase with trehalose [233,234] as well as combination of BHT with ascorbic acid [235] to cell types such as mononuclear cells, pancreatic islets, germ cells, spermatoozoa, dendritic cells, hepatocytes and hemopoietic cells (Table 1). Among these antioxidants, winter wheat lipocalins and peroxiredoxins obtained from wheat are especially notable. They were demonstrated to mollify cryopreservation-associated loss of attachment capacity of hepatocytes, as well as restoring the activity of CYP isoforms to the level similar from fresh, unfrozen murine hepatocytes [227,236]. Table 2 is a summary of the effect of antioxidants on non-reproductive cells.

Collectively, the different studies examined in this review indicated that the effectiveness of antioxidant supplement for cryopreservation very much depends on the cell type, organism as well as the specific antioxidant used. Table 1 provides a summary of the type of the antioxidant used, the cell type and organism as well as the effectiveness of the antioxidant based on the parameters measured.

**Moving forward**

Oxidative stress is inevitably generated in the cryopreservation process and has been widely cited as the causative factor for some of the cryoinjuries inflicted on the cell [163,165,187,237]. Therefore, administering antioxidants in an effort to counter these deleterious effects on cells during cryopreservation is a plausible solution. Indeed, the use of antioxidants has undoubtedly conferred protection to certain cell type by improving several cellular function parameters and general cryopreservation outcome in specific circumstances as those indicated in the sections above and in Tables 1 and 2. Although effective in some circumstances, antioxidants can be ineffective or even deleterious for some cells. Antioxidants consist of a broad class of substances and molecules with varying physio-chemical properties that dictate their specificity, localization and/or ROS-scavenging roles [238,239]. Mitochondria-targeted antioxidants, MitoTEMPO [214] and melatonin [240,241] are potent antioxidants that prevent oxidative stress-associated damages encountered during cryopreservation. Melatonin in particular, has performed unexpectedly well by exerting its ROS-ameliorating properties through its multi-faceted mechanisms [242]. While the administration of melatonin improved the generation and survival of somatic cell nuclear transferred (SCNT) murine embryos from vitrified oocytes, whether melatonin directly affects ROS or inhibits apoptosis remains to be elucidated [243]. As such, the use of antioxidant in different combinations for the different cell types for cryopreservation may prove to be more effective in countering cryopreservation-induced ROS damage.

There are evidences to indicate that the use of different antioxidants in combination could provide additive protective effect when compared with those administered individually (Tables 1 and 2). In reproductive cells, catalase and low concentration of SOD have been reported to have no effect on oocyte survivability and fertility when used alone. However, when the same dose of SOD was co-administered with catalase, significant improvement in oocyte survivability was observed [124]. For the case of sperm cryopreservation, supplementation of SOD alone has no effect on the general sperm parameters such as motility (total and progressive motility), viability and percentage of sperm with high MMP, while supplementation of catalase alone was beneficial only at high concentrations [244]. Notably, when catalase and SOD were used in combination, sperm parameters such as total motility was greatly improved as compared with individual use of them at the respective dose [244]. In another example, sperm cells frozen in SOD and catalase, or vitamins C and E led to significantly improved parameters such as reduced ROS [215] and increased
lateral head displacement of the sperm cells [245] whereas previously such parameters were not improved when the antioxidants were used individually [215,245].

Not all antioxidants used in combination yield additional benefits. For example, single administration of trehalose or catalase improved clonogenic parameters such as burst-forming unit erythroid and colony-forming unit granulocyte-monocyte in fetal liver hematopoietic cells and umbilical cord blood, respectively. However, when trehalose and catalase were administered in combination, no significant improvements in these parameters were observed [246]. Hence, use of different antioxidants in combination does not always imply additional improvement in cellular parameters.

Based on the studies examined in this review, it is notable that antioxidant supplement for cryopreservation can be effective. However, the effectiveness of the specific antioxidants depends on the cell type that undergoes the cryopreservation process. It is therefore important to consider supplementing cryopreservation media with specific antioxidants according to the specific species, cell type, quality and integrity of cells prior to cryopreservation. Additionally, several studies determined the efficacy of the antioxidants on cryopreservation by measuring functional parameters of the cryopreserved cells in an ex vivo setting. Given that many applications for cryopreservation are in the area of reproductive and regenerative medicine, future studies should be attempted to investigate the recovery and efficacy of the cryopreserved cells after transplantation in vivo to better understand the efficacy of the antioxidant used in this process.

Acknowledgments
The authors thank Drs Harmeet Singh, Yiu Wing Kam and Sharon Lim for proofreading the manuscript.

Author Contribution
J.S.L and W.S.D.K. wrote the manuscript, prepared the figures and tables. S.-X.T. supervised and edited the manuscript.

Funding
This work was supported by the Republic Polytechnic Research and Development Grant to S.-X.T.

Competing Interests
The authors declare that there are no competing interests associated with the manuscript.

Abbreviations
ATF6, activating transcription factor 6; BHT, butylated hydroxytoluene; CPA, cryoprotective agent; CYP, cytochrome P450; DHE, dihydroethidium; DMSO, dimethyl sulfoxide; DUOX, dual oxidase; ER, endoplasmic reticulum; ERO, ER oxidoreductase; ETC, electron transport chain; GSH, glutathione; H₂DCFDA, 2′,7′-dichlorodihydrofluorescein diacetate; H₂O₂, hydrogen peroxide; IRE1α, inositol-requiring enzyme 1α; L-NAME, NG-nitro-L-arginine methyl ester; MDA, malondialdehyde; MPP, mitochondrial membrane potential; mPTP, mitochondria permeability transition pore; NOS, nitric oxide synthase; NOX, NADPH oxidase; PBMC, peripheral blood mononuclear cell; PDI, protein disulfide isomerase; PERK, pancreatic eIF-2α kinase; ROS, reactive oxygen species; SAB, SH3 homology associated BTK binding protein 5; SOD, superoxide dismutase; TUDCA, tauroursodeoxycholic acid; UPR, unfolded protein response; XBP-1, X-box binding protein 1; 4-HNE, 4-Hydroxynonenal; γH2AX, H2A histone family member X.

References
1 Shu, Z., Heimfeld, S. and Gao, D. (2014) Hematopoietic SCT with cryopreserved grafts: adverse reactions after transplantation and cryoprotectant removal before infusion. Bone Marrow Transplant. 49, 469–476, https://doi.org/10.1038/bmt.2013.152
2 Lewis, J.K., Bischof, J.C., Braslavsky, I., Brockbank, K.G.M., Fahy, G.M., Fuller, B.J. et al. (2016) The grand challenges of organ banking: proceedings from the first global summit on complex tissue cryopreservation. Cryobiology 72, 169–182, https://doi.org/10.1016/j.cryobiol.2015.12.001
3 Varghese, A.C., du Plessis, S.S., Falcone, T. and Agarwal, A. (2008) Cryopreservation/transplantation of ovarian tissue and in vitro maturation of follicles and oocytes: challenges for fertility preservation. Reprod. Biol. Endocrinol. 6, 47
4 Moll, G., Alm, J.J., Davies, L.C., von Bahr, L., Heldring, N., Stenbeck Funke, L. et al. (2014) Do cryopreserved mesenchymal stromal cells display impaired immunomodulatory and therapeutic properties? Stem Cells 32, 2430–2442, https://doi.org/10.1002/stem.1729
5 Keith, Jr, S.C. (1913) Factors influencing the survival of bacteria at temperatures in the vicinity of the freezing point of water. Science 37, 877–879, https://doi.org/10.1126/science.37.962.877
6 Hollander, D.H. and Nell, E.E. (1954) Improved preservation of Treponema pallidum and other bacteria by freezing with glycerol. Appl. Environ. Microbiol. 2, 164–170
7 Macfadyen, A. and Rowland, S. (1902) On the suspension of life at low temperatures. *Ann. Bot. (Lond.) os-16*, 589–590, https://doi.org/10.1093/oxfordjournals.aob.a088893

8 Polge, C., Smith, A.U. and Parkes, A.S. (1949) Revival of spermatozoa after vitrification and dehydration at low temperatures. *Nature* 164, 666, https://doi.org/10.1038/164666a0

9 Polge, C. and Rowson, L.E.A. (1952) Fertilizing capacity of bull spermatozoa after freezing at −79°C. *Nature* 169, 626–627, https://doi.org/10.1038/169626b0

10 Lovelock, J.E. and Bishop, M.W.H. (1959) Prevention of freezing damage to living cells by dimethyl sulfoxide. *Nature* 183, 1394–1395, https://doi.org/10.1038/1831394a0

11 Ware, C.B., Nelson, A.M. and Blau, C.A. (2005) Controlled-rate freezing of human ES cells. *Biotechniques* 38, 879–883, https://doi.org/10.2144/05386ST01

12 Richards, M., Fong, C.Y., Tan, S., Chan, W.K. and Bongso, A. (2004) An efficient and safe xeno-free cryopreservation method for the storage of human embryonic stem cells. *Stem Cells* 22, 779–789, https://doi.org/10.1634/stemcells.22-5-779

13 Chinnadurai, R., Garcia, M.A., Sakurai, Y., Lam, W.A., Kirk, A.D., Galipeau, J. et al. (2011) Actin cytoskeletal disruption following cryopreservation alters the biodistribution of human mesenchymal stromal cells *in vivo*. *Stem Cell Rep.* 3, 60–72, https://doi.org/10.1016/j.stemcr.2014.05.003

14 Hattori, Y., Kato, H., Nitta, M. and Takamoto, S. (2001) Decrease of L-selectin expression on human CD34+ cells on freeze-thawing and rapid recovery with short-term incubation. *Exp. Hematol.* 29, 114–122, https://doi.org/10.1016/S0301-472X(00)00615-9

15 Katkov, I.I., Kim, M.S., Bajpai, R., Altman, Y.S., Mercola, M., Loring, J.F. et al. (2006) Cryopreservation by slow cooling with DMSO diminished production of Oct-4 pluripotency marker in human embryonic stem cells. *Cryobiology* 53, 194–205, https://doi.org/10.1016/j.cryobiol.2006.05.005

16 Wagh, V., Meganathan, K., Jagtap, S., Gaspar, J.A., Winkler, J., Spilkovskaya, D. et al. (2011) Effects of cryopreservation on the transcriptome of human embryonic stem cells after thawing and culturing. *Stem Cell Rev. Rep.* 7, 506–517, https://doi.org/10.1007/s12011-011-9230-1

17 Temple, M.D., Perrone, G.G. and Dawes, I.W. (2005) Complex cellular responses to reactive oxygen species. *Trends Cell Biol.* 15, 319–326, https://doi.org/10.1016/j.tcb.2004.05.003

18 Halliwell, B. (2006) Reactive species and antioxidants. Redox Biology is a fundamental theme of aerobic life. *Plant Physiol.* 141, 312–322, https://doi.org/10.1104/pp.106.077073

19 Halliwell, B. (2000) The antioxidant paradox. *Lancet* 355, 1179–1180, https://doi.org/10.1016/S0140-6736(00)02075-4

20 Schieber, M. and Chandel, N.S. (2014) ROS function in redox signaling and oxidative stress. *Curr. Biol.* 24, R453–R462, https://doi.org/10.1016/j.cub.2014.03.034

21 Finkel, T. and Holbrook, N.J. (2000) Oxidants, oxidative stress and the biology of ageing. *Nature* 406, 239–247, https://doi.org/10.1038/35041687

22 Biel, S., Faig, M., Tolahunase, M. and Dada, R. (2017) Oxidative stress and male infertility. *Nat. Rev. Urol.* 14, 470–485, https://doi.org/10.1038/nrurol.2017.69

23 von Zglinicki, T., Petrie, J. and Kirkwood, T.B.L. (2003) Telomere-driven replicative senescence is a stress response. *Nat. Biotechnol.* 21, 229–230, https://doi.org/10.1038/nbt0303-229b

24 Rizza, S., Cardaci, S., Montagna, C., Di Giacomo, G., De Zio, D., Bordi, M. et al. (2018) S-nitrosylation drives cell senescence and aging in mammals by controlling mitochondrial dynamics and mitophagy. *Proc. Natl. Acad. Sci. U.S.A.* 115, E3388–E3397, https://doi.org/10.1073/pnas.1722452115

25 Passos, J.F., Nelson, G., Wang, C., Richter, T., Simillion, C., Proctor, C.J. et al. (2010) Feedback between p21 and reactive oxygen production is necessary for cell senescence. *Mol. Syst. Biol.* 6, 347, https://doi.org/10.1038/msb.2010.5

26 Bigarella, C.L., Liang, R. and Ghaffari, S. (2014) Stem cells and the impact of ROS signalling. *Development* 141, 4206–4218, https://doi.org/10.1242/dev.107086

27 Nöžik Grayczyk, E., Huang, Y.C.T., Carriawaj, M.S. and Piantadosi, C.A. (2003) Bicarbonate-dependent superoxide release and pulmonary artery tone. *Am. J. Physiol. Heart Circ. Physiol.* 285, H2327–H2335, https://doi.org/10.1152/ajpheart.00507.2003

28 Mumbengegwi, D.R., Li, Q., Li, C., Bear, C.E. and Engelhardt, J.F. (2008) Evidence for a superoxide permeability pathway in endosomal membranes. *Mol. Cell. Biol.* 28, 3700–3712, https://doi.org/10.1128/MCB.02038-07

29 Fukai, T. and Uschio Fukai, M. (2011) Superoxide dismutases: role in redox signaling, vascular function, and diseases. *Antioxid. Redox Signal.* 15, 1583–1606

30 Chatterjee, S. and Gagnon, C. (2001) Production of reactive oxygen species by spermatozoa undergoing cooling, freezing, and thawing. *Mol. Reprod. Dev.* 59, 451–458, https://doi.org/10.1002/mrd.1052

31 Evangelista Vargas, S. and Santiani, A. (2017) Detection of intracellular reactive oxygen species (superoxide anion and hydrogen peroxide) and lipid peroxidation during cryopreservation of alpaca spermatozoa. *Reprod. Domest. Anim.* 52, 819–824, https://doi.org/10.1111/rda.12984

32 Honda, S., Weigel, A., Hjelmeland, L.M. and Handa, J.T. (2001) Induction of telomere shortening and replicative senescence by cryopreservation. *Biochem. Biophys. Res. Commun.* 282, 493–498, https://doi.org/10.1006/bbrc.2001.4585

33 Xu, X., Cowley, S., Flaim, C.J., James, W., Seymour, L. and Cui, Z. (2010) The roles of apoptotic pathways in the low recovery rate after cryopreservation of dissociated human embryonic stem cells. *Biotechnol. Prog.* 26, 827–837, https://doi.org/10.1021/bp100286d

34 Holt, C.B. (2003) Substances which inhibit ice nucleation: a review. *Cryoletters* 24, 269–274

35 Lee, H.H., Lee, H.J., Kim, H.J., Lee, J.H., Ko, Y., Kim, S.M. et al. (2015) Effects of antifreeze proteins on the vitrification of mouse oocytes: Comparison of three different antifreeze proteins. *Hum. Reprod.* 30, 2110–2119, https://doi.org/10.1093/humrep/dev170

36 Xu, X., Cowley, S., Flaim, C.J., James, W., Seymour, L.W. and Cui, Z. (2010) Enhancement of cell recovery for dissociated human embryonic stem cells after cryopreservation. *Biotechnol. Prog.* 26, 781–788, https://doi.org/10.1021/bp100358e

37 Dickinson, B.C. and Chang, C.J. (2011) Chemistry and biology of reactive oxygen species in signaling or stress responses. *Nat. Chem. Biol.* 7, 504–511, https://doi.org/10.1038/nchembio.607
Yoshikawa, T., Takahashi, S., Tanigawa, T., Naito, Y., Ichikawa, H., Takano, H. et al. (1991) Investigation into the reactivity between various amino acids and oxygen-derived free radicals by use of the ESR spin trapping method. *J. Clin. Biochem. Nutr.* **11**, 161–169, https://doi.org/10.3164/jcbn.11.161

Liu, F., Lai, S., Tong, H., Lakey, P.S.J., Shiraiva, M., Weiler, M.G. et al. (2017) Release of free amino acids upon oxidation of peptides and proteins by hydroxyl radicals. *Anal. Bioanal. Chem.* **409**, 2411–2420, https://doi.org/10.1007/s00216-017-088-y

Cadet, J., Delatoir, T., Douki, T., Gasparutto, D., Pouget, J.P., Ravanat, J.L. et al. (1999) Hydroxyl radicals and DNA base damage. *Mutat. Res.* **424**, 9–21, https://doi.org/10.1016/S0027-5107(99)00004-4

Gray, J.E., Starmer, J., Lin, V.S., Dickinson, B.C. and Magnuson, T. (2013) Mitochondrial hydrogen peroxide and defective cholesterol efflux prevent in vitro fertilization by cryopreserved inbred mouse sperm. *Biol. Reprod.* **89**, 1–12, https://doi.org/10.1095/biolreprod.113.109157

Imlay, J.A., Chin, S.M. and Linn, S. (1988) Toxic DNA damage by hydrogen peroxide through the Fenton reaction *in vivo* and *in vitro*. *Science* **240**, 640–642, https://doi.org/10.1126/science.2834821

Fridovich, I. (1998) Oxygen toxicity: a radical explanation. *J. Exp. Biol.* **201**, 1203–1209

Thomas, C., Mackey, M.M., Diaz, A.A. and Cox, D.P. (2009) Hydroxyl radical is produced via the Fenton reaction in submitochondrial particles under oxidative stress: implications for diseases associated with iron accumulation. *Redox Rep.* **14**, 102–108, https://doi.org/10.1016/j.redox.2013.05.012

Winterbourn, C.C. (2008) Reconciling the chemistry and biology of reactive oxygen species. *Nat. Chem. Biol.* **4**, 278–286, https://doi.org/10.1038/nchembio.85

Boveirs, A. and Chance, B. (1973) The mitochondrial generation of hydrogen peroxide. General properties and effect of hyperbaric oxygen. *Biochem. J.* **134**, 707–716, https://doi.org/10.1042/bj1340707

Adam Vizi, V. and Chinopoulos, C. (2006) Bioenergetics and the formation of mitochondrial reactive oxygen species. *FEBS Lett.* **580**, 335–344, https://doi.org/10.1016/j.febslet.2006.01.005

Mailloux, R.J. and Harper, M.E. (2012) Mitochondrial proctiy and ROS signaling: lessons from the uncoupling proteins. *Trends Endocrinol. Metab.* **23**, 451–458, https://doi.org/10.1016/j.tem.2012.04.004

Lambert, A.J. and Brand, M.D. (2004) Superoxide production by NADH:ubiquinone oxidoreductase (complex I) depends on the pH gradient across the mitochondrial inner membrane. *Biochem. J.* **382**, 511–517, https://doi.org/10.1042/BJ20040485

Murphy, M.P. (2009) How mitochondria produce reactive oxygen species. *Biochem. J.* **417**, 1–13, https://doi.org/10.1042/BJ20081386

Jones, A., Van Blerkom, J., Davis, P. and Toledo, A.A. (2004) Cryopreservation of metaphase II human oocytes affects mitochondrial membrane potential: Implications for developmental competence. *Hum. Reprod.* **19**, 1861–1866, https://doi.org/10.1093/humrep/deh313

Kadirel, G., Kumar, S. and Kumaresan, A. (2009) Lipid peroxidation, mitochondrial membrane potential and DNA integrity of spermatozoa in relation to intracellular reactive oxygen species in liquid and frozen-thawed buffalo semen. *Anim. Reprod. Sci.* **114**, 125–134, https://doi.org/10.1016/j.anireprosci.2008.10.002

Matsushita, T., Yagi, T., Hardin, J.A., Cragun, J.D., Crow, F.W., Bergen, III, H.R. et al. (2003) Apoptotic cell death and function of cryopreserved porcine hepatocytes in a bioartificial liver. *Cell Transplant.* **12**, 109–122, https://doi.org/10.3727/00000000003108746986

Starkov, A.A. and Fiskum, G. (2003) Regulation of brain mitochondrial H2O2 production by membrane potential and NAD(P)H redox state. *J. Neurochem.* **86**, 1101–1107, https://doi.org/10.1111/j.1471-4165.2003.01908.x

Korshunov, S.S., Skulachev, V.P. and Starkov, A.A. (1997) High protonic potential actuates a mechanism of production of reactive oxygen species in mitochondria. *FEBS Lett.* **416**, 15–18, https://doi.org/10.1016/S0014-5793(97)01159-9

Voytakova, T.V. and Reynolds, I.J. (2001) $\Delta \psi$-dependent and -independent production of reactive oxygen species by rat brain mitochondria. *J. Neurochem.* **79**, 266–277, https://doi.org/10.1046/j.1471-4165.2001.00548.x

Bolisetty, S. and Jaimes, E.A. (2013) Mitochondria and reactive oxygen species: physiology and pathophysiology. *Int. J. Mol. Sci.* **14**, 6306–6344, https://doi.org/10.3390/ijms14036306

Zorov, D.B., Filburn, C.R., Klotz, L.O., Zweier, J.L. and Sollott, S.J. (2000) Reactive oxygen species (ROS)-induced ROS release: a new phenomenon accompanying induction of the mitochondrial permeability transition in cardiac myocytes. *J. Exp. Med.* **192**, 1001–1014, https://doi.org/10.1084/jem.192.7.1001

Castro, L.S., Hamilton, T.R.S., Mendes, C.M., Richi, M., Barnabe, V.H., Visintin, J.A. et al. (2016) Sperm cryodamage occurs after rapid freezing phase: flow cytometry approach and antioxidant enzymes activity at different stages of cryopreservation. *J. Anim. Sci. Biotechnol.* **7**, 17, https://doi.org/10.1186/s40104-016-0076-x

Chen, S., Deng, X., Ma, K., Zhao, L., Huang, D., Li, Z. et al. (2018) Icariin improves the viability and function of cryopreserved human nucleus pulposus-derived mesenchymal stem cells. *Oxid. Med. Cell. Longev.* **2018**, 3459612, https://doi.org/10.1155/2018/3459612

Sohn, I.P., Ahn, H.J., Park, D.W., Gye, M.C., Jo, D.H., Kim, S.Y. et al. (2001) Amelioration of mitochondrial dysfunction and apoptosis of two-cell mouse embryos after freezing and thawing by the high frequency liquid nitrogen infusion. *Mol. Cells* **13**, 272–280
68 Slowinska, M., Liszewska, E., Judycka, S., Konopka, M. and Ciereszko, A. (2018) Mitochondrial membrane potential and reactive oxygen species in liquid stored and cryopreserved turkey (Meleagris gallopavo) spermatozoa. Poult. Sci. 97, 3709–3717, https://doi.org/10.3382/ps/pey020
69 Zee, Y.P., Holt, W.V., Allen, C.D., Nicolson, V., Burridge, M., Lisie, A. et al. (2007) Effects of cryopreservation on mitochondrial function and heterogeneity, lipid raft stability and phosphotyrosine translocation in koala (Phascolarctos cinereus) spermatozoa. Reprod. Fertil. Dev. 19, 850–860, https://doi.org/10.1071/RD07084
70 Demant, M., Trapphoff, T., Frohlich, T., Arnold, G.J. and Eichenlaub Ritter, U. (2012) Vitrification at the pre-antral stage transiently alters inner mitochondrial membrane potential but proteome of in vitro grown and matured mouse oocytes appears unaffected. Hum. Reprod. 27, 1096–1111, https://doi.org/10.1093/humrep/det45
71 Halestrap, A.P. (2009) What is the mitochondrial permeability transition pore? J. Mol. Cell Cardiol. 46, 821–831, https://doi.org/10.1016/j.yjmcc.2009.02.021
72 Petronilli, V., Penzo, D., Scorrano, L., Bernardi, P. and Di Lisa, F. (2001) The mitochondrial permeability transition, release of cytochrome c and cell death correlation with the duration of pore openings in situ. J. Biol. Chem. 276, 12030–12034, https://doi.org/10.1074/jbc.M010604200
73 Ortega Ferrusola, C., González Fernández, L., Salazar Sandoval, C., Macias Garcia, B., Rodríguez Martínez, H., Tapia, J.A. et al. (2010) Inhibition of the mitochondrial permeability transition pore reduces “apoptosis like” changes during cryopreservation of stallion spermatozoa. Theriogenology 74, 458–465, https://doi.org/10.1016/j.theriogenology.2010.02.029
74 Batandier, C., Leverve, X. and Fontaine, E. (2009) Opening of the mitochondrial permeability transition pore reduces “apoptosis like” changes during cryopreservation of stallion spermatozoa. Free Radic. Biol. Med. 99, 43–53, https://doi.org/10.1016/j.freeradbiomed.2016.07.026
75 Dong, Q., Tollner, T.L., Rodenburg, S.E., Hill, D.L. and Vandervoort, C.A. (2010) Antioxidants, oxyrase, and mitochondrial uncoupler 2.4-dinitrophenol improved post thaw survival of rhesus monkey sperm from ejaculates with low cryosurvival. Fertil. Steril. 94, 2359–2361, https://doi.org/10.1016/j.fertnstert.2010.04.017
76 Vercesi, A.E., Kowaltowski, A.J., Grijalba, M.T., Meinicke, A.R. and Castilho, R.F. (1997) The role of reactive oxygen species in mitochondrial permeability transition. Biochim. Biophys. Acta 1324, 130–135, https://doi.org/10.1016/S0005-2760(97)00047-8
77 Valle, V.G.R., Fagian, M.M., Parentoni, L.S., Meinicke, A.R. and Vercesi, A.E. (1993) The participation of reactive oxygen species and protein thiols in the mechanism of mitochondrial inner membrane permeabilization by calcium plus prooxidants. Arch. Biochem. Biophys. 307, 1–7, https://doi.org/10.1006/abbi.1993.1551
78 Zhao, N., Liu, X.J., Li, J.T., Zhang, L., Fu, Y., Zhang, Y.J. et al. (2015) Endoplasmic reticulum stress inhibition is a valid therapeutic strategy in vitrifying oocytes. Cryobiology 70, 48–52, https://doi.org/10.1016/j.cryobiol.2014.12.001
79 Barrera, N., dos Santos Neto, P.C., Cuadro, F., Bosoiasco, D., Mulet, A.P., Crispo, M. et al. (2018) Impact of delipidated estrous sheep serum supplementation on in vitro maturation, cytochrome and endoplasmic reticulum stress gene expression of sheep oocytes. PLoS ONE 13, e0198742, https://doi.org/10.1371/journal.pone.0198742
80 Brito, D.C., Brito, A.B., Scalercio, S.R.R.A., Percário, S., Miranda, M.S., Rocha, R.M. et al. (2014) Vitamin E-analog Trolox prevents endoplasmic reticulum stress in frozen-thawed ovarian tissue of capuchin monkey (Sapajus apella). Cell Tissue Res. 355, 471–480, https://doi.org/10.1007/s00441-013-1764-x
81 Lin, T., Lee, J.E., Shin, H.Y., Oqani, R., Kim, S.Y. and Jin, D.I. (2016) Supplement of tauroursodeoxycholic acid in vitrification solution improves postthaw survival of rhesus monkey sperm from ejaculates with low cryosurvival. Theriogenology 85, 575–580
82 Zee, Y.P., Holt, W.V., Allen, C.D., Nicolson, V., Burridge, M., Lisie, A. et al. (2007) Effects of cryopreservation on mitochondrial function and heterogeneity, lipid raft stability and phosphotyrosine translocation in koala (Phascolarctos cinereus) spermatozoa. Reprod. Fertil. Dev. 19, 850–860, https://doi.org/10.1071/RD07084
83 Demant, M., Trapphoff, T., Frohlich, T., Arnold, G.J. and Eichenlaub Ritter, U. (2012) Vitrification at the pre-antral stage transiently alters inner mitochondrial membrane potential but proteome of in vitro grown and matured mouse oocytes appears unaffected. Hum. Reprod. 27, 1096–1111, https://doi.org/10.1093/humrep/det45
84 Tu, B.P. and Weissman, J.S. (2004) Oxidative protein folding in eukaryotes. Science 304, 1096–1111, https://doi.org/10.1126/science.11047
85 Barrera, N., dos Santos Neto, P.C., Cuadro, F., Bosoiasco, D., Mulet, A.P., Crispo, M. et al. (2018) Impact of delipidated estrous sheep serum supplementation on in vitro maturation, cytochrome and endoplasmic reticulum stress gene expression of sheep oocytes. PLoS ONE 13, e0198742, https://doi.org/10.1371/journal.pone.0198742
86 Tu, B.P. and Weissman, J.S. (2004) Oxidative protein folding in eukaryotes. Science 304, 1096–1111, https://doi.org/10.1126/science.11047
87 Takahashi, S., Ando, A., Takagi, H. and Shima, J. (2009) Insufficiency of copper ion homeostasis causes freeze-thaw injury of yeast cells as revealed by indirect gene expression analysis. Appl. Environ. Microbiol. 75, 6705–6711, https://doi.org/10.1128/AEM.00905-09
88 Haynes, C.M., Titus, E.A. and Cooper, A.A. (2004) Degradation of misfolded proteins prevents ER-derived oxidative stress and cell death. Mol. Cell 15, 767–776, https://doi.org/10.1016/j.molcel.2004.08.025
89 Abraham, T., Pin, C.L. and Watson, A.J. (2012) Embryo collection induces transient activation of XBP1 arm of the ER stress response while embryo vitrification does not. Mol. Hum. Reprod. 18, 229–242, https://doi.org/10.1093/molehr/gar076
90 Kincaid, M.M. and Cooper, A.A. (2007) ERADicate ER stress or die trying. Antioxid. Redox Signal. 9, 2373–2387
91 Eletto, D., Chevet, E., Argon, Y. and Appenzeller Herzog, C. (2014) Redox controls UPR to control redox. J. Cell Sci. 127, 3649–3658, https://doi.org/10.1242/jcs.153643
92 Cuizzo, J.W. and Kaiser, C.A. (1999) Competition between glutathione and protein thiols for disulfide-bond formation. Nat. Cell Biol. 1, 130–135, https://doi.org/10.1038/11047
93 Tabas, I. and Ron, D. (2011) Integrating the mechanisms of apoptosis induced by endoplasmic reticulum stress. Nat. Cell Biol. 13, 184–190, https://doi.org/10.1038/nccbiot.2011-184
94 Haynes, C.M., Titus, E.A. and Cooper, A.A. (2004) Degradation of misfolded proteins prevents ER-derived oxidative stress and cell death. Mol. Cell 15, 767–776, https://doi.org/10.1016/j.molcel.2004.08.025
95 Kincaid, M.M. and Cooper, A.A. (2007) ERADicate ER stress or die trying. Antioxid. Redox Signal. 9, 2373–2387
154 Yousefian, I., Esmamverdi, M., Karamzadeh Dehaghi, A., Sabzian Melei, R., Zandi, M. and Zare Shahnai, A. (2018) Attenuation of cryopreservation-induced oxidative stress by antioxidant: impact of Coenzyme Q10 on the quality of post-thawed buck spermatozoa. Cryobiology 81, 88–93, https://doi.org/10.1016/j.cryobiol.2018.02.005

155 Isaac, A.V., Kumar, S., Nair, R., Urs, D.R., Sallam, S.R., Kalthur, G. et al. (2017) Supplementing zinc oxide nanoparticles to cryopreservation medium minimizes the freeze-thaw-induced damage to spermatozoa. Biochim. Biophys. Acta 1860, 310–318, https://doi.org/10.10111/j.1600-709k.2010.00843.x

156 Succu, S., Berlinguer, F., Pasciu, V., Satta, V., Leoni, G.G. and Naitana, S. (2011) Melatonin protects ram spermatozoa from cryopreservation injuries in a dose–dependent manner. J. Pineal Res. 50, 310–318, https://doi.org/10.1111/j.1600-709k.2010.00843.x

157 Bortolin, R.C., Garapoto, J., Vargas, A.R., da Silva Morrone, M., Kunzler, A., Henkin, B.S. et al. (2017) Effects of freeze-thaw and storage on enzymatic activities, protein oxidative damage, and immunocomponent of the blood, liver, and brain of rats. Biopreserv. Biobank. 15, 182–190, https://doi.org/10.1089/bio.2016.0023

158 Alotaibi, N.A.S., Slater, N.K.H. and Rahmoune, H. (2016) Salidroside as a novel protective agent to improve red blood cell cryopreservation. PLoS ONE 11, e0162748, https://doi.org/10.1371/journal.pone.0162748

159 Mostek, A., Dietrich, M.A., Slowińska, M. and Ciereszko, A. (2017) Cryopreservation of bull semen is associated with carbonylation of sperm proteins. Theriogenology 92, 95–102, https://doi.org/10.1016/j.theriogenology.2017.01.011

160 Tai, W.Y., Yang, Y.C., Lin, H.J., Huang, C.P., Cheng, Y.L., Chen, M.F. et al. (2010) Interplay between structure and fluidity of model lipid membranes under oxidative attack. J. Phys. Chem. B 114, 15642–15649, https://doi.org/10.1021/jp1014719

161 Yusupov, M., Wende, K., Kupsch, S., Neyts, E.C., Reuter, S. and Bogaerts, A. (2017) Effect of head group and lipid tail oxidation in the cell membrane revealed through integrated simulations and experiments. Sci. Rep. 7, 5761, https://doi.org/10.1038/s41598-017-06412-8

162 Whiteley, G.S.W., Fuller, B.J. and Hobbs, K.F. (1992) Deterioration of cold-stored tissue specimens due to lipid peroxidation: modulation by antioxidants at high subzero temperatures. Cryobiology 29, 668–673, https://doi.org/10.1016/0011-2240(92)90069-E

163 Ortega Ferrusola, C., González Fernández, L., Morrell, J.M., Salazar Sandoval, C., Macías García, B., Rodríguez Martínez, H. et al. (2009) Lipid peroxidation, assessed with BODIPY-C11, increases after cryopreservation of stallion spermatozoa, is stallion-dependent and is related to apoptotic-like changes. Reproduction 138, 55–63, https://doi.org/10.1530/REP-08-0484

164 Martin Muñoz, P., Ortega Ferrusola, C., Vizuete, G., Plaza Dávila, M., Rodríguez Martínez, H. and Peña, F.J. (2015) Depletion of intracellular thiols and increased production of 4-hydroxynonenal that occur during cryopreservation of stallion spermatozoa lead to caspase activation, loss of motility, and cell death. Biol. Reprod. 93, 143

165 Alvarez, J.G. and Storey, B.T. (1992) Evidence for increased lipid peroxidative damage and loss of superoxide dismutase activity as a mode of sublethal cryodamage to human sperm during cryopreservation. J. Androl. 13, 232–241

166 Schuffner, A., Morscheld, M. and Oehninger, S. (2001) Cryopreservation of fractionated, highly motile human spermatozoa: effect on membrane phosphatidylethanolamine externalization and lipid peroxidation. Hum. Reprod. 16, 2148–2153, https://doi.org/10.1093/humrep/16.10.2148

167 Puth, C.F., Berendsen, T.A., Bruinsma, B.G., Ozer, S., Luytje, M., Usta, O.B. et al. (2015) Polyethylene glycol protects primary hepatocytes during supercooling preservation. Cryobiology 71, 125–129, https://doi.org/10.1016/j.cryobiol.2015.04.010

168 Swami, D.S., Kumar, P., Malik, R.K., Saini, M., Kumar, D. and Jan, M.H. (2017) The cryoprotective effect of iodixanol in buffalo semen cryopreservation. Anim. Reprod. Sci. 198, 20–26, https://doi.org/10.1016/j.anireprosci.2017.01.012

169 Odani, M., Komatsu, Y., Oka, S. and Iwashashi, H. (2003) Screening of genes that respond to cryopreservation stress using yeast DNA microarray. Cryobiology 47, 155–164, https://doi.org/10.1006/cybi.2003.09.001

170 Park, J.I., Grant, C.M., Davies, M.J. and Dawes, I.W. (1998) The cytoplasmic Cu,Zn superoxide dismutase of Saccharomyces cerevisiae is required for resistance to freeze-thaw stress. Generation of free radicals during freezing and thawing. J. Biol. Chem. 273, 22921–22928, https://doi.org/10.1074/jbc.273.36.22921

171 Habihi, A., Farrokhi, N., Moreira da Silva, F., Bettencourt, B.F., Bruges Armas, J., Amid, F. et al. (2010) The effects of vitrification on gene expression in mature mouse oocytes by nested quantitative PCR. J. Assisted. Reprod. Genet. 27, 599–604, https://doi.org/10.1007/s10815-010-9453-0

172 Boonkulsom, D., Gal, A.B., Bodo, S., Gorhony, B., Kityanant, Y. and Dinnyes, A. (2006) Gene expression profiles and in vitro development following vitrification of pronuclear and 8-cell stage mouse embryos. Mol. Reprod. Dev. 73, 700–708, https://doi.org/10.1002/mrd.20450

173 Coppola, S. and Ghibelli, L. (2000) GSH extrusion and the mitochondrial pathway of apoptotic signalling. Biochem. Soc. Trans. 28, 56–61, https://doi.org/10.1042/bs0280056

174 Chen, G.Q., Ren, L., Zhang, D. and Shen, X.H. (2016) Glutathione improves survival of cryopreserved embryogenic calli of Agapanthus praecox subsp. orientalis. Acta Physiol. Plant. 38, 250, https://doi.org/10.1007/s10457-016-2271-y

175 Estrada, E., Rivera del Álamo, M.M., Rodríguez Gil, J.E. and Yeste, M. (2017) The addition of reduced glutathione to cryopreservation media induces changes in the structure of motile subpopulations of frozen-thawed boar sperm. Cryobiology 78, 56–64, https://doi.org/10.1016/j.cryobiol.2017.07.002

176 Moawad, A.R., Tan, S.L. and Taketo, T. (2017) Beneficial effects of glutathione supplementation during vitrification of mouse oocytes at the germinal vesicle stage on their preimplantation development following maturation and fertilization in vitro. Cryobiology 76, 98–103, https://doi.org/10.1016/j.cryobiol.2017.04.002

177 Vara, E., Arias Díaz, J., Villa, N., Hernández, J., Garca, C., Ortiz, P. et al. (1995) Beneficial effect of S-Adenosylmethionine during both cold storage and cryopreservation of isolated hepatocytes. Cryobiology 32, 422–427, https://doi.org/10.1006/cryo.1995.1041

178 Ha, S.J., Kim, B.G., Lee, Y.A., Kim, Y.H., Kim, B.J., Jung, S.E. et al. (2016) Effect of antioxidants and apoptosis inhibitors on cryopreservation of murine germ cells enriched for spermatogonial stem cells. PLoS ONE 11, e0161372, https://doi.org/10.1371/journal.pone.0161372

179 Hwang, C., Sinskey, A. and Lodish, H. (1992) Oxidized redox state of glutathione in the endoplasmic reticulum. Science 257, 1496–1502, https://doi.org/10.1126/science.1523409
208 Gottlieb, E., Armour, S.M., Harris, M.H. and Thompson, C.B. (2003) Mitochondrial membrane potential regulates matrix configuration and cytochrome c release during apoptosis. Cell Death Differ. 10, 709–717, https://doi.org/10.1038/sj.cdd.4401231

209 Scarlett, J.L., Sheard, P.W., Hughes, G., Ledgewood, E.C., Ku, H.H. and Murphy, M.P. (2000) Changes in mitochondrial membrane potential during staurosporine-induced apoptosis in Jurkat cells. FEBS Lett. 475, 267–272, https://doi.org/10.1016/S0014-5793(00)01681-1

210 Gottlieb, E., Vander Heiden, M.G. and Thompson, C.B. (2000) Bcl-xL prevents the initial decrease in mitochondrial membrane potential and subsequent reactive oxygen species production during tumor necrosis factor alpha-induced apoptosis. Mol. Cell. Biol. 20, 5680–5689, https://doi.org/10.1128/MCB.20.15.5680-5689.2000

211 Zamzami, N., Marchetti, P., Castedo, M., Decaudin, D., Macho, A., Hirsch, T. et al. (1995) Sequential reduction of mitochondrial transmembrane potential and generation of reactive oxygen species in early programmed cell death. J. Exp. Med. 182, 367–377, https://doi.org/10.1084/jem.182.2.367

212 Hardikar, A.A., Risbud, M.V., Remacle, C., Reusens, B., Hoet, J.J. and Bhonde, R.R. (2001) Islet cryopreservation: improved recovery following taurine pretreatment. FEBS Lett. 506, 315–320, https://doi.org/10.1016/S0014-5793(01)03006-0

213 Peña, F.J., Johansson, A., Wallgren, M. and Rodríguez Martinez, H. (2003) Antioxidant supplementation in vitro improves boar sperm motility and mitochondrial membrane potential after cryopreservation of different fractions of the ejaculate. Anim. Reprod. Sci. 78, 85–98, https://doi.org/10.1016/S0378-4320(03)00049-6

214 Lu, X., Zhang, Y., Bai, H., Liu, J., Li, J. and Wu, B. (2018) Mitochondria-targeted antioxidant MitoTEMPO improves the post-thaw sperm quality. Cryobiology 80, 26–29, https://doi.org/10.1016/j.cryobiol.2017.12.009

215 Figuerola, E., Farias, J.G., Lee Estevez, M., Valdebenito, I., Risopatrón, J., Magnotti, C. et al. (2018) Sperm cryopreservation with supplementation of α-tocopherol and ascorbic acid in freezing media increase sperm function and fertility rate in Atlantic salmon (Salmo salar). Aquaculture 493, 1–8, https://doi.org/10.1016/j.aquaculture.2018.04.046

216 Silva, E.C.B., Cajuéiro, J.F.P., Silva, S.V., Soares, P.C. and Guerra, M.M.P. (2012) Effect of antioxidants resveratrol and quercetin on in vitro evaluation of Atlantic salmon sperm. Theriogenology 77, 1722–1726, https://doi.org/10.1016/j.theriogenology.2011.11.023

217 Zhang, L., Xue, Y., Yao, J., Yan, L.Y., Jin, X.H., Zhu, X.H. et al. (2016) L-proline: a highly effective cryoprotectant for mice oocyte vitrification. Sci. Rep. 6, 26326, https://doi.org/10.1038/srep26326

218 Carneiro, J.A.M., Canisso, I.F., Bandeira, R.S., Scheeren, V.F.C., Freitas Dell’Aqua, C.P., Alvarenga, M.A. et al. (2018) Effects of coenzyme Q10 on cryopreservation of stallions classified as having good or bad semen freezing ability. Anim. Reprod. Sci. 192, 107–118, https://doi.org/10.1016/j.anireprosci.2018.02.020

219 Garcez, M.E., dos Santos Branco, C., Lara, L.V., Pasqualotto, F.F. and Salvador, M. (2010) Effects of resveratrol supplementation on semen cryopreservation of medium human semen. Fertil. Steril. 94, 2118–2121, https://doi.org/10.1016/j.fertnstert.2010.01.058

220 Kotdawala, A.P., Kumar, S., Sialani, S.R., Thankachan, P., Govindraj, K., Kumar, P. et al. (2012) Addition of Zinc to human ejaculate prior to cryopreservation prevents freeze-thaw-induced DNA damage and preserves sperm function. J. Assist. Reprod. Genet. 29, 1447–1453, https://doi.org/10.1007/s10815-012-9994-8

221 Limaye, L.S. (1997) Bone marrow cryopreservation: improved recovery due to bioantioxidant additives in the freezing solution. Stem Cells 15, 353–358, https://doi.org/10.1002/stem.150353

222 Hardikar, A.A., Risbud, M.V., Remacle, C., Rouaix, B., Hoet, J.J. and Bhonde, R.R. (2001) Islet cryopreservation: improved recovery following taurine pretreatment. Cell Transplant. 10, 247–253, https://doi.org/10.3727/000000001783986756

223 De Loever, P., Fuller, B.J. and De Loecker, W. (1991) The effects of cryopreservation on protein synthesis and membrane transport in isolated rat liver mitochondria. Cryobiology 28, 445–453, https://doi.org/10.1016/0011-2240(91)90053-Q

224 Manning Fox, J.E., Lyon, J., Dai, X.O., Wright, R.C., Hayward, J., van de Bunt, M. et al. (2015) Human islet function following 20 years of cryogenic biobanking. Diabetologia 58, 1503–1512, https://doi.org/10.1007/s00125-015-3598-4

225 Mukherjee, N., Chen, Z., Sambanis, A. and Song, Y. (2005) Effects of cryopreservation on cell viability and insulin secretion in a model tissue-engineered pancreatic substitute (TEPS). Cell Transplant. 14, 449–456, https://doi.org/10.3727/000000005783982882

226 Jitaruch, S., Dhawan, A., Hughes, R.D., Filippi, C., Lehec, S.C., Glover, L. et al. (2017) Cryopreservation of hepatocyte microbeads for clinical transplantation. Cell Transplant. 26, 1341–1354, https://doi.org/10.1080/000000005783982882

227 Jitaruch, S., Dhawan, A., Hughes, R.D., Filippi, C., Lehec, S.C., Glover, L. et al. (2017) Cryopreservation of hepatocyte microbeads for clinical transplantation. Cell Transplant. 26, 1341–1354, https://doi.org/10.1080/000000005783982882

228 Grondin, M., Hamel, F., Averill-Bates, D. and Sarhany, F. (2009) Wheat proteins improve cryopreservation of rat hepatocytes. Biootechnol. Bioeng. 103, 582–591, https://doi.org/10.1002/bit.22270

229 Mugishima, H., Harada, K., Chin, M., Suzuki, T., Takagi, K., Hayakawa, S. et al. (1999) Effects of long-term cryopreservation on hematopoietic progenitor cells in umbilical cord blood. Bone Marrow Transplant. 23, 395–396, https://doi.org/10.1038/sj.bmt.1701580

230 Othmani, A.E., Rouam, S., Abbad, A., Erraoui, M., Harriba, S., Boukind, H. et al. (2019) Cryopreservation impacts cell functionality of long term expanded adipose-derived stem cells. J. Stem Cell Res. Ther. 9, 445

231 Zhang, X.B., Li, K., Yao, K.H., Tsang, K.S., Fok, T.F., Li, C.K. et al. (2003) Trehalose ameliorates the cryopreservation of cord blood in a preclinical system and increases the recovery of CFUs, long-term culture-initiating cells, and nonobese diabetic-SCID repopulating cells. Transfusion 43, 265–272, https://doi.org/10.1046/j.1537-2995.2003.00301.x

232 Xue, W.J., Luo, X.H., Li, Y., Liu, H.B., Tian, X.H., Feng, X.S. et al. (2011) Effects of astaxanthin on cultured islets after cryopreservation in rats. Transplant. Proc. 43, 3908–3912, https://doi.org/10.1016/j.transproceed.2011.10.039

233 Sasnoor, L.M., Kale, V.P. and Limaye, L.S. (2005) Prevention of apoptosis as a possible mechanism behind improved cryoprotection of hematopoietic cells by catalase and trehalose. Transplantation 80, 1251–1260, https://doi.org/10.1097/01.tp.0000169028.01327.90
234 Sasnoor, L.M., Kale, V.P. and Limaye, L.S. (2005) A combination of catalase and trehalose as additives to conventional freezing medium results in improved cryoprotection of human hematopoietic cells with reference to in vitro migration and adhesion properties. Transfusion 45, 622–633, https://doi.org/10.1111/j.0041-1132.2005.04288.x

235 Fujita, R., Hui, T., Chelly, M. and Demetriou, A.A. (2005) The effect of antioxidants and a caspase inhibitor on cryopreserved rat hepatocytes. Cell Transplant. 14, 391–396, https://doi.org/10.3727/000000005783982981

236 Chow-shi-yé, M., Grondin, M., Averill Bates, D.A. and Ouellet, F. (2016) Plant protein 2-Cys peroxiredoxin TaBAS1 alleviates oxidative and nitrosative stresses incurred during cryopreservation of mammalian cells. Biotechnol. Bioeng. 113, 1511–1521, https://doi.org/10.1002/bit.25921

237 Tatone, C., Di Emidio, G., Vento, M., Cirimmina, R. and Artini, P.G. (2010) Cryopreservation and oxidative stress in reproductive cells. Gynecol. Endocrinol. 26, 563–567, https://doi.org/10.3109/09513591003686395

238 Nimse, S.B. and Pal, D. (2015) Free radicals, natural antioxidants, and their reaction mechanisms. RSC Adv. 5, 27986–28006, https://doi.org/10.1039/C4RA13315C

239 Halliwell, B. (1996) Antioxidants: the basics—what they are and how to evaluate them. Adv. Pharmacol. 38, 3–20, https://doi.org/10.1016/S1054-3589(08)60976-X

240 Zhu, Z., Li, R., Lv, Y. and Zeng, W. (2019) Melatonin protects rabbit spermatozoa from cryo-damage via decreasing oxidative stress. Cryobiology 88, 1–8, https://doi.org/10.1016/j.cryobiol.2019.04.009

241 Appiah, M.O., He, B., Lu, W. and Wang, J. (2019) Antioxidative effect of melatonin on cryopreserved chicken semen. Cryobiology 89, 90–95, https://doi.org/10.1016/j.cryobiol.2019.05.001

242 Reiter, R.J., Mayo, J.C., Tan, D.X., Sanz, R.M., Alatorre Jimenez, M. and Qin, L. (2016) Melatonin as an antioxidant: under promises but over delivers. J. Pineal Res. 61, 253–278, https://doi.org/10.1111/jpi.12360

243 Lee, A.R., Hong, K., Choi, S.H., Park, C., Park, J.K., Lee, J.I. et al. (2019) Anti-apoptotic regulation contributes to the successful nuclear reprogramming using cryopreserved oocytes. Stem Cell Rep. 12, 545–556, https://doi.org/10.1016/j.stemcr.2019.01.019

244 Trzcińska, M. and Bryla, M. (2015) Apoptotic-like changes of boar spermatozoa in freezing media supplemented with different antioxidants. J. Vet. Sci. 18, 473–480

245 Roca, J., Rodriguez, M.J., Gill, M.A., Carvajal, G., Garcia, E.M., Cuello, C. et al. (2005) Survival and in vitro fertility of boar spermatozoa frozen in the presence of superoxide dismutase and/or catalase. J. Androl. 26, 15–24

246 Limaye, L.S. and Kale, V.P. (2001) Cryopreservation of human hematopoietic cells with membrane stabilizers and bioantioxidants as additives in the conventional freezing medium. J. Hematother. Stem Cell Res. 10, 709–718

247 Kim, G.A., Lee, S.T., Ahn, J.Y., Park, J.H. and Lim, J.M. (2010) Improved viability of freeze-thawed embryonic stem cells after exposure to glutathione. Fertil. Steril. 94, 2409–2412, https://doi.org/10.1016/j.fertnstert.2010.01.073