Cryopreservation of Human Stem Cells for Clinical Application: A Review

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Summary
Stem cells have been used in a clinical setting for many years. Haematopoietic stem cells have been used for the treatment of both haematological and non-haematological disease; while more recently mesenchymal stem cells derived from bone marrow have been the subject of both laboratory and early clinical studies. Whilst these cells show both multipotency and expansion potential, they nonetheless do not form stable cell lines in culture which is likely to limit the breadth of their application in the field of regenerative medicine. Human embryonic stem cells are pluripotent cells, capable of forming stable cell lines which retain the capacity to differentiate into cells from all three germ layers. This makes them of significant importance in both regenerative medicine and toxicology. Induced pluripotent stem (iPS) cells may also provide a similar breadth of utility without some of the confounding ethical issues surrounding embryonic stem cells. An essential prerequisite to the commercial and clinical application of stem cells are suitable cryopreservation protocols for long-term storage. Whilst effective methods for cryopreservation and storage have been developed for haematopoietic and mesenchymal stem cells, embryonic cells and iPS cells have proved more refractory. This paper reviews the current state of cryopreservation as it pertains to stem cells and in particular the embryonic and iPS cell.
Introduction

The application of stem cells to treat human disease is by no means new. Bone marrow was first transplanted in the 1950s [1, 2], although the first major breakthrough came in 1968 with two reports of successful allogeneic transplantation in two patients: one with an X-linked lymphopenic immune deficiency and the other with Wiskott-Aldrich syndrome [3, 4]. More recently, in 1988, the first successful transplant was performed using umbilical cord blood on a child with Fanconi’s anaemia [5]. Whether sourced from bone marrow, umbilical cord blood or, predominantly today from mobilised peripheral blood, the haematopoietic stem cell has developed into the most common source of cells for cell-based therapies for both haematological and non-haematological disease [6, 7]. A recent world-wide retrospective survey of haematopoietic stem cell transplantation (HSCT) from 1,327 participating centres in 71 countries reported over 51,000 first HSCTs (43% allogeneic) for 2006 [8], while the most recent activity survey from the European Group for Blood and Marrow Transplantation (EBMT), reporting for 2008, showed a total of over 30,000 HSCTs of which 90% were first transplants [9]. Whilst the main indications were predominantly for leukaemias and lymphomas, over 12% were for solid tumours and non-malignant disorders.

Mesenchymal Stem Cells

A second, separate non-haematopoietic stem cell population with multi-lineage potential, residing in the bone marrow, was identified by Friedenstein et al. in 1976 [10]. These stromal cells, variously known as mesenchymal stem cells, marrow stromal stem cells, or mesenchymal stromal cells (MSCs), are capable of differentiation to give rise to bone (osteocytes), cartilage (chondrocytes) and fat (adipocytes) [11] when induced ex vivo. In addition to bone marrow, these cells have been reported in an increasing range of tissues including peripheral blood [12], umbilical cord blood [13], placenta [14], amniotic fluid and membrane [15, 16], dental pulp [17], deciduous teeth [18] and adipose tissue [19]. The incidence of these cells in tissue is extremely low, ranging from around 0.00003% of nucleated cells in cord blood to 0.001–0.01% of nucleated cells in marrow, though this decreases with age [20]. Adipose tissue has been shown to have a higher proportion of MSCs (approximately 2% in the stromal vascular fraction) with a differentiation capacity that extends beyond the mesenchymal lineages [21]. A rich source of MSCs is the perivascular compartment surrounding the vessels of the umbilical cord (with a reported frequency of 1 in 300 of the nucleated cell population) [22]. These have also demonstrated multilineage potential in clonal studies; differentiating to five mesenchymal lineages: bone, cartilage, fat, muscle, and fibrous tissue [23]. The plasticity of MSCs has been well documented [24], and successful differentiation to a wide variety of lineages has been demonstrated including those above and also cardiomyocytes [25], hepatocytes [26], neurons [27] and endothelium [28, 29] though not without question [30, 31]. Recently, a multipotent stem cell (termed a multi-lineage-differentiating, stress-enduring or ‘Muse’ cell) capable of giving rise to all three germ layers and expressing a set of genes associated with pluripotency, including Oct 3/4, Sox 2 and Nanog, has been isolated and clonally selected from bone marrow stromal cells [32]. A similar pluripotent stem cell type has also been isolated from human umbilical cord blood [33]. This, termed an unrestricted somatic stem cell, could also be differentiated to all three germ layers but did not express the major pluripotency factors. However it did display an epigenetic signature that suggested a ‘poised’ epigenetic state for the pluripotency genes, thus preserving the cells’ pluripotent potential [34]. These cell types may help to explain the reported ability of MSCs to differentiate to cells of all three germ layers.

Unlike the haematopoietic stem cells from bone marrow and cord blood, plastic-adherent MSCs can be expanded in culture without the loss of differentiation capacity. However these cells show a finite life span of anywhere between 15 and 50 population doublings [24, 35] before successive passaging reduces the proliferative capacity and multi-lineage differentiation potential [36, 37]. Moreover, and unlike haematopoietic stem cells which express common surface markers (e.g. CD34 and CD133), MSCs lack a unique identifying phenotypic marker and are characterised by a set of minimum criteria including the presence/absence of certain specific markers and their in vitro differentiation capacity in response to specific stimuli [38]. Whilst their expansion capability permits the development of a number of MSC-based therapies [39, 40], this finite lifespan, passage-related loss of differentiation capacity, and lack of a specific marker or markers has implications in terms of quality control and batch variation when generating stem cells for commercial and clinical application.

Embryonic Stem Cells

Pluripotent human embryonic stem (hES) cells, unlike MSCs, have, for all practical purposes, an unlimited capacity for self-renewal and in culture maintain their pluripotent capacity to differentiate into cell types from all three germ layers, as demonstrated by their ability to form teratomas in SCID mice. Though first isolated from surplus in vitro fertilised blastocysts in 1994 [41], it was not until 1998 that the first stable hES cell line was established by Thomson et al. [42]. The capacity of these cells to undergo virtually infinite expansion and asymmetric cell division with differentiation into cells of ectodermal, endodermal and mesodermal origin was confirmed some 2 years later [43]. Since then, hundreds of stem cell lines have been derived world-wide from blastocysts of fresh and cryopreserved supernumerary embryos as well as from morula [44], single blastomeres [45], arrested embryos and embryos discarded after pre-implantation genetic diagnosis [46, 47].
Phenotypically and in contrast to MSCs, hES cells express a set of surface and intracellular markers which, while not exclusive to embryonic stem cells, nevertheless characterise these cells. This canonical set of markers includes the stage-specific embryonic antigens SSEA-3 and SSEA-4, TRA-1-60, TRA-1-81, CD9 and CD133 (also expressed by other stem cell populations). Thy-1 (CD90), MHC class 1 and the intracellular transcription factor Oct 3/4 [48]. Undifferentiated cells also show high levels of alkaline phosphatase and telomerase activity and generally maintain a normal karyotype over extended periods in culture though there have been some reports of culture adaptation generally, but not exclusively, at high passage involving karyotypic and genotypic changes [49, 50].

**Induced Pluripotent Stem Cells**

A very recent development, with potentially a profound significance for clinical therapy has been the generation of induced pluripotent stem (iPS) cells from somatic cells [51–53]. The reprogramming of human somatic cells to an embryonic stem cell-like pluripotent state, through the forced expression of key transcription factors, was first demonstrated by Takahashi et al. in 2007 [54] concurrently with two other groups [55, 56]. Almost immediately it was shown that such cells could be generated from patient-specific cells for a wide variety of disease states [57] and from a wide variety of somatic cell types [58]. The generation and use of iPS cells particularly for autologous stem cell therapy poses fewer ethical problems when compared to the derivation and use of embryonic stem cells. However, whilst these cells have been shown to be embryonic-stem cell-like, there are small but potentially significant differences in their gene expression patterns [59]; the functional significance and effect on clinical utility of which remain uncertain. The methods of generation too pose some concern for their clinical application: the use of lentiviral and retroviral vectors as well as of oncogenes (Myc and Klf4) together with the low efficiency of reprogramming techniques may compromise their growth and developmental characteristics as well as their clinical utility [60]. Differences between iPS and hES cells in their relative ability to undergo directed differentiation and a recent report that iPS cells derived from different adult tissues varied substantially in their teratoma-forming properties [61] all argue for caution and continued research with hES cells if the full potential of stem cell therapies is to be realised.

**Stem Cells for Clinical Therapy**

The ability of stem cells from whatever source, but particularly embryonic stem cells, to produce a theoretically unlimited supply of normal, differentiated cells has focussed attention on the potential importance of these cells in both toxicology and drug discovery [62, 63], tissue engineering [64, 65] as well as gene and cellular therapy [66, 67] for a wide range of human diseases including Parkinson’s and other neurodegenerative diseases [68–70], diabetes [71, 72] and cardiac and vascular therapy [73, 74]. In a recent editorial in *Regenerative Medicine* [75], Mason and Manzotti, using a definition for a regenerative medicine as ‘that which replaces or regenerates human cells, tissue or organs to restore normal function’, argue that the cell therapy sector (both public and commercial) is likely to impact on a huge diversity of medical specialties and applications ranging from conventional cell, tissue and organ transplantation, through transient cell therapies that disrupt or reduce natural disease progression, to tissue engineering and bioaesthetic cell-based treatments. Respected commentators on the potential of the regenerative medicine field have plotted the development of regenerative medicine therapies since the early 2000s and have anticipated a substantial growth in this area based on progress so far [77]. By 2007 over a million patients had been treated, and European licensing authorities were seeing a sharp increase in the number of applications in the pipeline. From a commercial perspective, this positive benefit to human health has a current world-wide market for regenerative medicine of between USD 2 and 5 billion, with US sales of commercially available stem cell therapies alone valued at USD 15.2 million in 2007 and USD 16.5 million in the first 2 quarters of 2008. The stem cell market, including therapies, cord blood banking and drug development tools, has been projected to achieve annual growth of 29%, resulting in sales of more than USD 11 billion by 2020. World-wide, there are currently over 800 clinical trials listed on the US Clinical Trials website which are recruiting to studies in which the search terms ‘mesenchymal stem cell’ or ‘adult stem cell’ are used and ‘stem cell transplant’ appears in the intervention [78]. MSCs are being used clinically in trials as diverse as cirrhosis, left ventricular dysfunction, and graft-versus-host disease [79, 80], while trials of a neural stem cell line for the treatment of stroke patients and oligodendrocyte progenitor cells derived from hES cells for the treatment of spinal injury have just been announced [80, 81]. However, in order to provide safe, quality-controlled cells for both autologous and allogeneic clinical therapy, regardless of the cell source, the principles of good manufacturing practice (GMP) must be applied.

**GMP in the Production of Cells for Clinical Therapies**

GMP, a quality assurance system used by pharmaceutical manufacturers, is used in the production of stem cells for clinical application. GMP regulations, first promoted by the US Food and Drug Administration, and implemented in Europe as EU Directive 2003/94/EC [83], underpin the EU Tissue and Cells Directives and the Advanced Therapy Medicinal Products Regulations which pertain to the production of haematopoietic, MSCs, hES and iPS cells for clinical therapy within most of Europe. Within the USA these guidelines have been incorporated into the Good Tissue Practice requirements
under the Code of Federal Regulations [84, 85]. Internationally, harmonisation of regulation is taking place through the International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH) of which the guidelines ICHQ7 ‘Good Manufacturing Practice’, ICHQ9, ‘Quality Risk Management’ and ICHQ10 ‘Pharmaceutical Quality Systems’ are particularly relevant [86].

The purpose of GMP is to ensure that these cell-based products are safe, pure and effective, by seeking to control all aspects of the production process from donor selection and harvesting/derivation of the cells, through enrichment/expansion to cell preservation, storage and distribution. It requires traceability throughout the production chain and beyond (through systems for product recall and adverse event reporting) and is predicated on appropriate risk assessment and risk mitigation.

At its most simple, as with haematopoietic stem cells, processing involves mainly volume reduction, preservation and storage. Systems have been developed over the years to manage the risk associated with this form of minimal processing: mainly the avoidance of microbiological contamination and the maintenance of the CD34+ cell population [87–89], and standards have been laid down to ensure that the cells are clinically acceptable [90–91]. However, in cell production processes that involve expansion and banking of cells over periods of weeks and months, the maintenance of conditions that ensure safety and efficacy is particularly onerous, and it is a common misconception that laboratory practices need only be recapitulated by the simple expedient of transferring the process to a cleanroom facility for the cells to be suitable for clinical application. Whilst such facilities are important in the prevention of environmental contamination, all aspects of the process from derivation/isolation of the cells through their maintenance in culture to the preservation, storage and transportation of the cells must be risk assessed for their potential to contribute hazards both microbiologically and non-microbiologically that may affect the safety and efficacy of the cells.

Many of the risks entailed in the banking of stem cells, particularly as they relate to MSCs, hES and iPSC cells, are similar to those for any cell culture process. It is well known that cell lines maintained in serial culture are susceptible to genetic variation. The formation of frozen banks of homogeneous cell aliquots can therefore capture a single desired phenotype allowing researchers to share source material containing functionally identical cells that maintain the desired characteristics; making possible direct comparison between studies performed by different research groups or by single researchers over time.

Additionally, the risk of contamination increase when cell lines are maintained for long periods of time in culture. Cell culture medium will readily support the growth of bacteria and fungi and the routine use of antibiotics to suppress growth may have adverse effects on the cells. While such contamination is often readily identified by the turbidity of the culture medium, more subtle forms of contamination such as mycoplasma introduced from the environment; viruses introduced via components of the culture medium and even contamination by other cell lines, are less readily detected and will affect the properties of the cells. Thus, returning periodically to a well characterised, contamination-free, frozen stock of cells reduces the risk of contamination and increases the validity of any results generated from the cell line.

It is not the intention here to review cell culture procedures, the use of xenogeneic components for derivation/isolation/cell expansion, or the risks associated with these; this has been done elsewhere [84, 89, 92–94]. The purpose of this paper is to review cryopreservation, low-temperature storage and transportation of stem cells, particularly as these apply to the embryonic and iPSC cell, where there are significant problems both in preservation and compliance with the requirements of GMP.

**Cryopreservation of Haematopoietic Stem Cells**

This routine procedure generally involves slow cooling in the presence of a cryoprotectant to avoid the damaging effects of intracellular ice formation. The cryoprotectant in popular use is dimethyl sulphoxide (DMSO), and the use of a controlled rate freezing technique at 1 to 2 °C/min and rapid thawing is considered standard [95]. Passive cooling devices that employ mechanical refrigerators, generally at −80 °C, to cool the cells (so-called dump-freezing) generate cooling rates similar to those adopted in controlled rate freezing. Generally, the outcome from such protocols has been comparable to controlled rate freezing [96–98] though the lack of a freezing record such as that produced by controlled rate cooling machines is problematic from a regulatory perspective.

Whilst the current cryopreservation protocols are clinically effective, questions still remain as to whether or not they are optimal. This is of particular concern in the banking and therapeutic application of cord blood for allogeneic use, where harvested volumes are small and the total number of CD34+ cells/kg body mass is crucial to eventual engraftment in the recipient. Here a number of studies have been undertaken in order to replace the largely empirical approach to developing an optimised protocol with a methodological one that takes into account the sequence of damaging events that occur during the freezing and thawing process [99, 100]. Such studies have shown that, whilst cooling rates of 1–2.5 °C/min are probably optimal for DMSO concentrations of 5–10%, recovery is likely to be improved if the osmotic damage that occurs through the introduction and removal of the cryoprotectant is tempered by the application of slow addition/elution protocols [101].

DMSO is known to be toxic to tissues and cells, with toxicity being time-, temperature- and concentration-dependent.
Toxicity varies from cell type to cell type, and the accepted practice has been to introduce the cryoprotectant at low temperatures (+4 °C) for as short a period as is considered practical. However, our studies and others have shown that the temperature-dependent toxicity, at the concentrations used in haematopoietic cells cryopreservation, is unlikely to be of practical importance even at ambient temperatures [101, 102].

Adverse reactions in patients receiving cryopreserved haematopoietic stem cells due to the accepted practice of re-infusing the thawed cells without removal of the DMSO are well documented [95], and washing procedures generally based on that originally developed by Rubenstein et al. [103] are in routine use. This protocol, though only employing a two-step elution and washout procedure for DMSO, is carried out in the presence of Dextran. This acts as an osmotic buffer preventing damaging osmotic transients during the removal of the cryoprotectant. The washing procedure also helps to remove granulocyte debris which will also help to reduce adverse reactions in the recipient. Other procedures involving automated cell washing devices have been developed [95, 100].

Reduced concentrations of DMSO have also been employed as a means to alleviate adverse reactions where washing procedures were not employed. There seems little adverse effect on cell recovery or engraftment in reducing DMSO concentration to 5% at optimal cooling rates [101, 104], and concentrations as low as 2% have been successfully employed [95]. Alternative cryoprotectants such as hydroxethyl starch and trehalose, either in combination with DMSO [105, 106] or alone [107], have also been shown to be effective in cryopreserving haematopoietic cells, but only so far in laboratory studies.

**Cryopreservation of Mesenchymal Stem Cells**

MSCs from bone marrow and cord blood are cryopreservable by the methods used to cryopreserve their haematopoietic analogues, and slow freezing protocols utilising DMSO as cryoprotectant are in use by many groups [108]. Recent studies by Liu et al. [109] have confirmed that 10% DMSO and slow cooling / rapid warming does not affect the viability or differentiation potential of adipose-derived MSCs. Adult stem cells derived from human dental pulp also showed high post-thaw recovery and trilineage differentiation potential after slow cooling in 1–1.5 mol/l DMSO (~7.5–10%). Furthermore, DMSO was shown to be superior to both ethylene glycol and propylene glycol under these conditions [110].

DMSO is known to have an effect on the epigenetic profile of, and induce differentiation in, murine stem cells [111, 112]. This, together with the known adverse reactions when employed as a cryoprotectant for haematopoietic cell therapy, plus its cellular toxicity, have led to attempts to cryopreserve MSCs with other cryoprotectants; either on their own or in combination with DMSO at lowered concentrations. Many of the alternative cryoprotectant formulations have also attempted to remove animal serum from the cryoprotectant solution; both to reduce cost and to improve clinical utility through reducing the likelihood of zoonotic infections in the absence of terminal sterilisation procedures. This has led to substitution with human serum or human serum albumin. However, this too is costly and introduces at least the potential for transmission of human pathogens.

The use of polyvinylpyrrolidone (PVP), an extracellular cryoprotectant, has been investigated as an alternative to cryopreservation with DMSO and foetal calf serum (FCS) [113]. Cellular recovery and differentiation capacity were studied after equilibration in a number of different cryopreservation media followed by 'dump' freezing to ~80 °C and storage in liquid nitrogen. Recovery of cells cryopreserved in 10% PVP with human serum was comparable with (though slightly lower than) cells cryopreserved in DMSO with animal serum. A similar study utilising methylcellulose either alone or in conjunction with reduced levels of DMSO indicated that human serum could replace FCS in standard DMSO mixtures without affecting the recovery of cells and that 1% methylcellulose produced comparable results with DMSO concentrations as low as 2% when an annexin V apoptosis assay was used to analyses cells 24 h post-thaw. Cells also maintained their adipogenic and osteogenic potential [114]. Lui et al. [115] have also adopted a reduced DMSO approach in a recent study to produce a well-defined and xeno-free cryopreservation media for cell therapy with bone marrow-derived MSCs. They replaced FCS with combinations of polyethylene glycol (PEG) and trehalose, with DMSO concentration ranging from 2.5 to 7.5%. The recovery, metabolic activity, proliferative capacity and differentiation potential were measured against the standard slow cooling protocol of DMSO and 10% FCS. Results replacing FCS with PEG in reduced concentrations of DMSO were comparable with those of the control, but this was achieved only in the presence of 2% bovine serum albumin. Moreover, cryopreservation in 10% DMSO with 90% FCS was more effective in their hands than any other combination. Trehalose was ineffective, in contrast to studies on haematopoietic cells derived from cord blood, bone marrow or mobilised peripheral blood where its effectiveness in combination with reduced DMSO concentration has been demonstrated [116, 117]. Nevertheless, these laboratory studies indicate that there is a potential for the development of xeno-free cryoprotectant solutions utilising lowered concentrations of DMSO and slow cooling.

**Cryopreservation of Embryonic and Induced Pluripotent Stem Cells**

Whilst both haematopoietic cells and MSCs have proved amenable to cryopreservation by conventional slow cooling protocols, their pluripotent counterparts have been shown to
be more refractory. In early studies, using a standard slow cooling protocol, Reubinoff et al. [118] reported 16% recovery after freezing and thawing (as measured by the number of colonies recovered 2 weeks after thawing). The colonies recovered were undersized compared to typical hES cell colonies and showed a significant degree of differentiation. Zhou et al. [119] also reported similar results with only slightly higher recovery (approximately 23%, based on the number of colonies recovered at day 9, post-thaw). Again, the rate of growth of the surviving colonies was reduced, while the level of differentiation in the surviving colonies was higher than that in a comparable, non-cryopreserved group. Both studies utilised 10% DMSO as the cryoprotectant and slow cooling at around 1 °C/min. A study by Ha et al. [120] compared glycerol and ethylene glycol (EG) to DMSO, either on their own or in combination with DMSO. The results indicated that, even with the most favourable combination (5% DMSO combined with 10% EG in 50% FCS) recovery was limited to just 30%. Results from studies by Heng et al. [121] confirmed this apparent failure of embryonic stem cells to survive conventional cryopreservation protocols.

The problem of effective cryopreservation is exacerbated by the ‘co-operative’ nature of these cells. Unlike adult stem cells, which grow in culture as adherent monolayers of un-associated cells, human (and animal) embryonic stem cells (and iPSCs) will only normally grow in the undifferentiated state as colonies of tightly associated, adherent cells (typically containing between 3 × 10^4 and 5 × 10^4 cells per hES cell colony) in co-culture with mitotically inactivated mouse or human fibroblasts as a feeder layer. During cell culture, mammalian cells are sub-cultured by some form of enzymatic dissociation, followed by agitation to remove the cells from the underlying substratum. Dissociation is completed through to a single cell suspension. It is in this form that the cells are passed or treated with cryoprotectant and cryopreserved.

Mouse embryonic stem cell lines, though colonial in nature, are routinely sub-cultured in this way, and may be successfully cryopreserved as an undifferentiated, single cell suspension [122]. In contrast, hES cells (and primate embryonic stem cells) have not been amenable to passing or cryopreserving as single cell suspensions. Primate stem cells show an extremely low plating efficiency if dissociated to single cells [123] while re-plating hES cells as a single cell suspension tends to lead to differentiation or cell death. Optimum cell cluster size for cryopreservation has been reported to be in the order of 100–500 cells [124]; though experimental evidence is sparse.

The practical consequence of this, together with the poor results obtained through conventional cryopreservation, led the derivation groups to seek alternative methods of cryopreservation that avoided the damaging effects of slow cooling. This led to the adoption of vitrification as the method of choice for the preservation of embryonic stem cell lines. Whilst this poses some problems for the efficient banking of cell lines for research purposes, there are additional consequences in terms of the application of GMP to this process for cells intended for clinical applications.

**Cryopreservation by Vitrification**

During conventional slow cooling, ice forms as a response to the reduction in temperature below the freezing point of the system. The consequences of this are an increase in the solute concentration as water is converted to ice. Both ice and high solute concentration are damaging to cells. Cryoprotectants such as DMSO mitigate these damaging effects by reducing the amount of ice formed at any given sub-zero temperature, thereby limiting the damaging rise in solute concentration. Vitrification approaches the problems associated with ice crystallisation in a different manner. Here cryoprotectants are used in concentrations that prevent ice formation entirely.

Vitrification is the solidification of a liquid without crystallisation and the growth of ice. This is achieved when solutes in the system are sufficiently concentrated, or the system cooled sufficiently rapidly, that the increased viscosity inhibits nucleation and prevents the growth of ice. As cooling continues, viscosity continues to increase until all molecular motion is (for all practical purposes) halted and the solution becomes a glass. In this condition, the system displays the properties of a solid but retains the molecular structure of a liquid [125].

Vitrification may be achieved by increasing the concentration of cryoprotectants to levels sufficient to avoid ice formation whatever the cooling rate. However, this requires the introduction of much higher concentrations of cryoprotectants than are usually tolerated by cells. Chemical toxicity is time-, temperature- and concentration-dependent, and this approach to vitrification has generally required both the careful formulation of multimolar cryoprotectant mixtures, to help lower toxicity and their introduction in a stepwise fashion at increasingly lower temperatures. However osmotic damage is increased by reduction in temperature, and thus any protocol for vitrification using this approach is often a compromise between inflicting chemical toxicity or osmotic damage on the cell. Such an approach has been termed equilibrium vitrification. An alternative approach is non-equilibrium vitrification. This relies on employing extremely high cooling rates in conjunction with lower concentrations of cryoprotectant in order to prevent ice nucleating. However this produces a metastable state that can lead to devitrification (and ice crystallisation) on re-warming. Though this poses serious risks of cell damage, this approach has been successfully applied to many cell and tissues, including embryos, cord blood and amnion-derived MSCs [126–128].

The adoption of this method of vitrification as the predominant method of cryopreservation for hES cells is largely due to comparative studies by three groups, two of which indicated rates of recovery of undifferentiated colonies of >75% for vitrified hES cells compared to ~5% after slow cooling [118, 119, 129]. The vitrification protocols reported in these
studies were very similar, based as they were on that developed for bovine ova and embryos [130] and modified by Reubinoff et al. [118] for application to hES cells. The method has also been described in detail with minor modifications elsewhere [131].

In essence, this protocol requires the stepwise exposure of hESC colony fragments to two vitrification solutions of increasing cryoprotectant concentration, the common components of which are DMSO and EG. The composition of the vehicle solution varies, with differences in sucrose concentration, the presence or absence of serum and the buffer used. Colony fragments are exposed to the two vitrification solutions sequentially. Exposure to the vitrification solutions is brief (60 s and 25 s respectively at either room temperature or 37 °C). No studies have so far been reported to determine the permeability of the (or the colony fragments) to either cryoprotectant, or the intrinsic toxicity of these components to hES cells; though studies are underway.

In non-equilibrium vitrification, the higher the cooling rate attained, the lower can be the total cryoprotectant concentration needed and vice versa. Using mixtures of cryoprotectants helps to reduce the intrinsic toxicity of each, and the method published by Reubinoff et al. [118] utilised 20% DMSO, 20% EG and 0.5 mol/l sucrose. To achieve vitrification using these solutions, very rapid cooling rates were required, though the critical cooling velocity needed to achieve vitrification has not been determined for this solution. Applying the OPS method (for open pulled straw), the high cooling rate was achieved by the direct immersion in liquid nitrogen of finely drawn capillaries holding ultra-small volumes (~20 μl or less) of the vitrification solution containing the colony fragments. Straws were then generally transferred under liquid nitrogen (to avoid devitrification) to liquid-phase nitrogen storage [130, 132].

To avoid ice crystallisation during thawing the straws were re-warmed as rapidly as possible by direct immersion of the tip of the loaded straw into pre-warmed culture medium containing sucrose. Once thawed the colony fragments were then expelled into this medium and transferred stepwise through cryoprotectant wash-out solutions, containing decreasing concentration of sucrose as the osmotic buffer, until they were plated into culture medium. An alternative method with direct exposure to growth medium without stepwise elution of the cryoprotectant has also been used with no noticeable detrimental effects [131].

However, vitrification by this technique is technically challenging, and there are anecdotal reports that recovery rates are not as high in practice as those reported in the literature. Even the original paper by Reubinoff et al. [118] reported recovery of <30% undifferentiated colonies (a figure not incomparable with those published for slow cooling). The method also suffers from two inherent problems that affect its application in a commercial, GMP setting: scale-up for the production of large banks of cells and the direct exposure of the cells to liquid nitrogen [127].

Vitrification and Scale-Up

The limitations on sample volume, necessary to achieve the required cooling rates, and the need to vitrify the embryonic stem cells as small clumps of stem cells at best permits the production of only laboratory-scale cell banks by the OPS method. This process is time-consuming, operator-dependent and, with the short exposure times to cryoprotectant, prone to both failure and inconsistency in the final product.

Attempts have been made to address this, at least on a laboratory scale. Heng et al. [133] proposed a design for a culture plate with detachable wells in which whole adherent colonies could be vitrified. Harvesting the hES cell clumps after colony dissociation and passage through a nylon cell strainer, followed by exposure to vitrification solutions and vitrification by direct immersion in liquid nitrogen whilst still retained on the strainer, has been employed by Li et al. [134] for the ‘bulk’ vitrification of embryonic stem cells. Using this method, the equivalent of a 35 mm culture dish (approximately 100–150 cell clumps) could be transferred to a single cell strainer, (equivalent to 10–20 straws) in a procedure taking no more than 5 min. When compared to hES cells recovered by the OPS method, no significant differences were detected in either the rate of re-attachment or in the degree of differentiation (measured at day 7) nor in the pluripotency of the surviving cells. A refinement of this technique, using customised cryovials fitted with stainless steel mesh, produced similar results [135]. Whilst both methods allow for an increase in the quantity of hES cells preserved at any one time, neither method is suitable for automation and commercial scale-up nor do they overcome the potential contamination problems associated with direct exposure to liquid nitrogen.

Vitrification and Contamination

The transmission of infections through in vitro fertilisation routes has been the subject of a number of reviews [136, 137], and much therein is relevant to embryonic stem cell lines. Moreover, transmission of infection associated with direct exposure to liquid nitrogen has been the source of a number of reports from both laboratory and clinical studies [138, 139]. Though methods and equipment exists and have been validated for the sterile filtration of liquid nitrogen [140], its applicability to cell banking and sterility at the point of use is questionable. Thus current vitrification techniques utilising open straws, followed by their storage under liquid nitrogen is unlikely to be acceptable by regulatory agencies for ‘clinical grade’ pluripotent stem cell lines.

Methods to separate the vitrification solution from the oxygen have been developed. The simplest of these is the sealed straw used in embryo and gamete cryopreservation. Successful vitrification of hES cell colony fragments has been demonstrated by Richards et al. [129] using sterile 250 μl embryo straws which could be sealed using a conventional heat sealer. Such straws have been approved for use by the US Food and
Drug administration [137]. The results were comparable to the OPS method when comparable volumes (i.e. 20 μl) of hES cell-containing vitrification solution were used. Whilst cooling rates for direct immersion of these straws were not measured, a study by Vajta et al. [130] comparing open, manually pulled straws with conventional 250 μl plastic straws indicated that, for similar volumes of vitrification solution, an order of magnitude difference in cooling rate was obtained (~22,000 °C/min vs. ~2,500 °C/min). Whether or not the colony fragments are truly vitrified in the Richards et al. study [129], cooling (and re-warming) is nevertheless sufficiently rapid to afford good rates of recovery.

Whilst the sealed straw prevents contact with potentially contaminating liquid nitrogen during cooling, it does not prevent the possible contamination of the external surface of the straw during handling and storage and its transmission to the thawing and recovery solution. This potential contamination route has been recognised in tissue and haematopoietic stem cell banking and is combated by the use of a secondary container (so called double-bagging). A straw-in-straw method, eliminating this potential route of contamination has been successfully employed for mouse embryos [141] and more recently mouse neurospheres [142]. This technique is likely to cause a further reduction in the cooling (and subsequent rewarming) rate possibly by a further order of magnitude [127]. Thus the cooling rate is unlikely to be sufficiently rapid to sustain vitrification of hES cells in the absence of changes to the composition of the vitrification solutions, and its potential as a closed system for vitrification of human hES and iPS cells has yet to be investigated.

The use of cryovials for vitrification has also been explored. Nishigaki et al. [143] successfully cryopreserved primary embryonic stem and hES cells in a vitrification solution without DMSO. They compared a number of vitrification solutions employing various concentrations of DMSO, EG and PEG with knock-out serum (KSR) replacement in both DMEM and Eurocollins vehicle solutions. Plunge cooling rates for 200 μl volumes were measured at ~125 °C/min. Differential scanning calorimetry (DSC) measurements on the solutions (containing 40% EG, 10% PEG and 20% KSR) indicated that the solutions would vitrify at this cooling rate. A re-attachment index was used to assess post-warming recovery after 1 day in culture, and they obtained recovery rates of around 20% by this method with surviving colonies expressing typical pluripotency markers and the ability to form teratomas.

Another approach to the problem of direct contact between cells and cryogen has been the development of solid surface vitrification. In essence this separates the cryogenic fluid from the samples by using the cryogen to cool a sterile solid surface [144] onto which the microdroplets of vitrification solution containing the cells or tissue are dropped. This solid surface [144] onto which the microdroplets of vitrification fluid from the samples by using the cryogen to cool a sterile surface vitrification. In essence this separates the cryogenic fluid from the samples by using the cryogen to cool a sterile solid surface [144] onto which the microdroplets of vitrification solution containing the cells or tissue are dropped. This method has now been commercialised and used to vitrify murine ovarian tissue and porcine blastocysts [145, 146].

Whilst these methods address the problems associated with potential contamination via the cryogen and other methods have increased the through-put of material that can be vitrified, no method combines both of these requirements nor lends itself to scale-up or automation. At best the sealed straw and straw-in-straw methods may provide a means, acceptable to the regulatory authorities, for small-scale cell banking of seed stocks of stems cells for therapeutic application. For these and for other logistical reasons, the cryopreservation method for clinical applications is most likely to involve conventional slow cooling methodologies despite their apparent problems.

**Cryopreservation by Conventional Slow Cooling**

As already discussed the early studies on cryopreserving hES cells lines using slow cooling techniques and DMSO with varying concentrations of serum were discouraging [118–121, 129]. These studies all employed protocols previously effective for murine embryonic stem (mES) cell lines which in turn had been adapted from conventional slow cooling protocols for cultures cells [147]. Unlike hES cells, when applied to mES cells, recovery after thawing was typically in excess of 90% with surviving cells showing low rates of differentiation and high rates of proliferation whilst still retaining their pluripotent capacity.

These early comparative studies made little attempt to control or investigate any of the cryobiological variables associated with damage during slow cooling: cooling rate, cryoprotectant concentration, osmotic effects associated with the introduction and removal of the cryoprotectant, or, in a system where the cellular material is frozen as an agglomerate of cells requiring maintenance of cell-to-cell contact, ice nucleation. The problems were (and continue to be) exacerbated by the necessity to apply suitable assays to measure not only immediate post-thaw recovery in cell clumps of varying and undefined size, but also the ability of these agglomerates to re-attach, proliferate and maintain pluripotency for extended periods in culture and the n-points at which these assays are applied [148]. Comparison of studies on both vitrification and/or slow cooling is therefore often difficult, and the importance placed on evidence of pluripotency and karyotypic stability as indicators of the effectiveness of the cryopreservation procedure is perhaps overemphasised. Such tests are generally carried out after a number of days or weeks in culture where the effects of sub-lethal injury will have been diluted out [149].

Though no studies have adopted a methodological approach similar to that employed for haematopoietic [99, 104] or mES cells [150], a number of more recent studies have attempted to investigate some of the cryobiological variables. These studies have indicated that control of the cryobiological variables can significantly improve the recovery of embryonic stem cells preserved by slow cooling. In the first of these studies, Ware et al. [151] investigated the effect of seeding on the
survival of a range of hES cell lines after thawing. Survival was calculated as combined colony number / colony size, relative to a pre-freeze control. Clusters were frozen by controlled-rate cooling in 10% DMSO at cooling rates between 0.3 and 3 °C/min. High survival (~80%) was obtained at cooling rates below 1.8 °C/min, but only if the samples were seeded. The effect of seeding temperature on survival was not fully investigated; however, seeding at –7 and –10°C produced similar results. Yang et al. [152] have reported similar results with seeding at –10 °C and a cooling rate of 0.5 °C/min.

Valbuena et al. [153] investigated the effect of stepwise addition of 2 mol/l DMSO in a four-step addition process at room temperature on hES cells. These were then subjected either to passive cooling at –80 °C overnight or two-step cooling without seeding. Though immediate post-thaw survival was low (~10%), it was comparable in the cell lines studied to cells vitrified by the OPS and sealed straw methods. More recent and encouraging studies have emphasised the beneficial effects of stepwise equilibration of the cryoprotectant and seeding during slow cooling. Li et al. [154] obtained re-attachment and recovery rates of over 50% using programmed cooling (1 °C/min) with seeding at –7 °C after single-step addition of the cryoprotectant (10% DMSO) with comparatively low survival in the group cryopreserved without seeding. A similar study [155] including two-step equilibration of the cryoprotectant also demonstrated improved recovery after seeding at –7 °C (>50%) as well as highlighting another possible confounding factor: variability in the cell line response to slow cooling injury. In this study two genetically modified embryonic stem cells lines were separated into four groups depending on method of cryoprotectant addition and presence or absence of a seeding event in the slow cooling programme. In all groups one of the two lines consistently exhibited an approximately 50% lower level of recovery than the other. In the group cryopreserved by single-step addition of cryoprotectant and no seeding, results were comparable to those obtained in many of the other studies (~5%). At this level of recovery it is difficult to effectively demonstrate differences that could be attributable to the cell line itself and, since the difference is likely to be attributable to the cells osmotic response to cryoprotectant exposure, would not have been visible in many of the early studies.

In a multifactorial study using Rhesus Macaque embryonic stem cells, Baran and Ware [156] demonstrated firstly the superiority of DMSO over EG or a mixture of EG and DMSO, secondly an optimum cooling rate of around 1 °C/min (compared to 0.3 °C/min or plunge freezing in 10% DMSO), and, lastly, an effect of both warming rate and post-thaw exposure to DMSO at 37 °C (equivalent to thermal runaway) on the survival after freezing. This study did not investigate the effect of stepwise equilibration but did again demonstrate that controlled-rate cooling with the induction of ice nucleation could increase post-thaw survival dramatically, from <22 to >90%, at least in this primate stem cell line.

Mechanisms of Slow Cooling Injury

The basic principles and damaging events that can underlie slow cooling injury have been well documented [157]. Cryoprotectant toxicity, osmotic damage during exposure to and removal of cryoprotectants, and so-called solution effect injury from the high solute concentrations to which the cells are exposed during freezing are just some of the damaging events that may occur. No systematic studies have yet been undertaken in the context of hES cells; however there is evidence of the protective effect of seeding to control ice formation within the system.

During slow cooling of a cell suspension, ice nucleation will generally be initiated in the extracellular compartment with cells inhabiting the solute channels that form between the growing ice crystals. Where nucleation has occurred within a cell, intracellular ice will form but this event will not affect neighbouring cells. In a cluster of cells such an event may have more profound consequences. Cell-to-cell propagation of intracellular ice has been demonstrated in cultured cell monolayers [158] and cell strands from insect salivary glands [159]. It has been postulated that this occurs via gap junctions, and there is experimental evidence to support this. Acker et al. [160] compared ice propagation in MDCK (gap junction forming) and V-79W (non-gap junction-forming) cell monolayers and observed a significant difference in the formation and propagation of intracellular ice in confluent monolayers with and without gap junctions. Cooling rates employed in this study were in the order of 25 °C/min; however an increase in the incidence of cell damage, attributed to the damaging effect of ice nucleation in cell monolayers compared to single cell suspensions, has been reported for corneal keratinocytes at 1 °C/min [161].

The presence of functional gap junctions has been demonstrated in hES cells [162, 163], and gap junction communication has been implicated in a number of cellular processes including cell proliferation, differentiation and apoptosis [164, 165], leading to speculation that these structures contribute to paracrine signalling between hES cells and are essential for survival and control of the undifferentiated state [42, 43, 124]; control that is disrupted if gap junctions are disrupted by pas- saging as single cell suspensions or cryopreservation. The observation that intracellular ice propagates more readily between cells with gap junctions under certain conditions may at least in part explain the poor recovery of hES cells after slow cooling. Damage to the hESC clusters caused by intercellular ice propagation, either by random nucleation events within the cluster and propagation through the gap junctions [166] or from surface-catalysed nucleation at its periphery [167] followed by cell-to-cell propagation, could lead to disruption of the cell cluster affecting cell proliferation, differentiation and apoptosis on thawing [159]. Methods that attempt to control the nucleation event, such as seeding, may therefore act to initiate ice formation outside of the cellular (or cell cluster) compartment.
Modification to the environment in which the cells experience the freezing process also appears to modify the extent of cellular injury. Ji et al. [168] compared whole colonies frozen in suspension with colonies frozen whilst still adherent to the culture surface with only a thin layer of cryoprotectant covering the cell layer. Survival in both cases was low (<10%). However when the adherent cells were embedded within a layer of Matrigel (a laminin-based basement membrane matrix for anchorage-dependent cells), immediate post-thaw viability was increased. Subsequent re-plating gave a higher incidence of re-attachment and a lower incidence of differentiation compared to colonies frozen in suspension or frozen adherent but un-encapsulated, perhaps through preferential formation of ice in the encapsulating matrix.

Apoptosis in response to low temperature exposure is well documented [169], and the involvement of apoptotic cell death in cryopreservation injury has been reported in a wide variety of cell types including haematopoietic stem cells [170]. Heng et al. [171] postulated the involvement of apoptosis in early cell death in hES cells cryopreserved by slow cooling after observing that cell viability immediately post-thaw (as measured by trypan blue dye exclusion) was initially very high but showed a gradual reduction with time in culture at 37°C. Moreover this loss of viability could be reversibly slowed by a reduction in the temperature at which the cells were held post-thaw [172], indicating an apoptotic mechanism for cell death rather than an unregulated necrotic process. The role of cryopreservation in activating both the extrinsic and intrinsic pathways of apoptosis was recently investigated by Xu et al. [173]. They examined the effect of DMSO exposure and slow cooling on the production of reactive oxygen species (ROS), p53 levels, levels of the initiators of the apoptotic cascade (caspases 8 and 9) and cytoskeletal F-actin (which can contribute to the induction of apoptosis in response to environmental changes). Propidium iodide (PI) / annexin V FITC staining by FACS 2 h post-thaw was used to assess initial viability and early apoptosis. PI-positive cells accounted for approximately 20% of the cell population with another 30% exhibiting early-stage apoptosis (PI-negative, annexin V-positive). Exposure to DMSO had no effect on the level of hydrogen peroxide or superoxide anion generation, however there were significant increases in these following freezing and thawing. They concluded that elevation of ROS led to the activation and translocation of p53 as strong expression of this protein was seen in the nucleus of post-thawed cells. This in turn led to activation of caspase-9 which was also significantly increased after thawing. Caspase-8 activity also showed a similar increase post-thaw, indicating the possible activation of the extrinsic pathway.

Caspase inhibitors significantly reduced caspase activity, but this did not appear to affect an improvement in post-thaw recovery, in contrast to other studies [174]. Addition of an inhibitor of Bax (a pro-apoptotic protein regulating mitochondrial membrane permeability) did however improve post-thaw recovery though not to the same extent as either the Rho kinase (ROCK) inhibitor Y-27632 or ROCK plus a p53 inhibitor (pifithrin-μ). The inhibition by ROCK of both caspase-8 and caspase-9 activity led them to the conclusion that cryopreservation activated both the extrinsic and intrinsic apoptotic pathways in hES cells; though the particular sensitivity of hES cells to apoptosis as compared to mES cells remains unanswered, even though both show similar sensitivities to ROS (which requires the addition of an antioxidant to their routine culture medium).

**Rho-Associated Kinase (ROCK) Inhibitors and the Survival of Stem Cells**

Rho kinases are proteins which have been shown to play a significant role in an array of cellular processes, including adhesion, proliferation, differentiation and apoptosis depending on the cell type [175]. The ROCK inhibitors Y-27632 and fasudil were first shown to markedly reduce dissociation-induced apoptosis in hES cells by Watanabi et al. [176] when these cells were dissociated to single cell suspensions prior to passaging. However, their exact mode of action in hES cells is a matter of some debate. It has been suggested that instead of blocking apoptotic pathways, ROCK inhibitors counteract anoikis (detachment-induced apoptosis) or enhance cell-cell interactions through modulation of gap junctions thereby increasing the adhesive properties of the cells and enhancing post-dissociation aggregation [177, 178]. Whatever the mode of action, ROCK inhibitors have been shown to improve post-thaw survival in a number of studies. Li et al. [179] demonstrated that Y-27632 added to the post-thaw culture medium for 24 h post-thaw increased hES cell survival rate approximately 10-fold, with colony growth rates similar to unfrozen controls.

Studies by Martin-Ibanez et al. [180] replicated this initial finding. Addition of the inhibitor Y-27632 to the cryopreservation medium alone did not provide any statistically significant improvement compared to cells frozen in the absence of the inhibitor. However, they too demonstrated a beneficial effect of its inclusion post-thaw. Furthermore, addition to both the preservation solution and post-thaw culture medium further improved survival; with hES cell colonies exhibiting low levels of differentiation. They also measured Oct-4 levels and the canonical pluripotency markers. Cells displayed pluripotency markers at day 1 post-thaw and Oct-4 levels, as measured by RT-PCR, were comparable to non-frozen controls at all post-thaw time points.

The effect of ROCK inhibitor Y-27632 has now been shown to have similar effects on the recovery from cryopreservation of both adult stem cells and bone marrow-derived MSCs as well as human iPS cells in both feeder-associated and feeder-free conditions [181–184] and protocols for the cryopreservation of dissociated hES cells have recently been published [185, 186].

**Other Additives and Alternative Cryoprotectants**

Alternatives to DMSO as the cryoprotectant of choice for stem cells have been driven in non-embryonic stem cells
largely by the adverse reactions caused in patients. In hES cells the motivation has been the known effect of this solvent on inducing differentiation [187]. However the minimum concentration (0.125%) and exposure time (days) required for it to exert an effect are likely to be of importance only in toxicological studies utilising hES cells, and not in the hES cell culture or cell banking.

Cryopreservation using the non-permeating disaccharide trehalose has been compared with slow cooling in 10% DMSO and serum [168]. Trehalose does not readily penetrate cells, with only low concentrations (<0.5%) being absorbed by passive diffusion or active endocytotic mechanisms. Cells were exposed to the trehalose loading medium for 24 h prior to exposure to the standard cryoprotectant mixture (10% DMSO and serum). There was a small but statistically significant increase in post-thaw survival of adherent colonies compared to DMSO serum without trehalose but only where the serum concentration in the cryoprotectant medium exceeded 50%. Wu et al. [188] also utilised trehalose. They showed a beneficial effect of trehalose in the elution solution, suggesting its mode of action as an osmotic buffer, and in the freezing medium (without a pre-incubation), suggesting that the trehalose was acting as an extracellular cryoprotectant.

Polyampholytes (polyelectrolytes bearing both cationic and anionic repeat group) such as poly-L-lysine have been used successfully to cryopreserve rat MSCs [189] and may offer an alternative to DMSO. They offer the advantage of also being antifreeze proteins that may help to control or reduce ice crystallisation. Sericin (a protein derived from the silkworm) has been shown to accelerate cell proliferation in hybridoma cell lines [190] and improve the attachment of cryopreserved hepatocytes, when used as a replacement for serum in a DMSO-based cryoprotectant with maltose [191].

Antioxidants in the form of β-mercaptoethanol are routinely present in the culture medium of hES cells but the overproduction of ROS during cryopreservation has led to the addition of antioxidants to the freezing solution in an attempt to reduce damage. Adding glutathione to the cryoprotectant and the post-thaw recovery solution has been shown to improve the survival of embryonic stem cells following cryopreservation [192].

Commercial Cryoprotectant Solutions

The drive towards clinical applications has led to the re-appraisal of both the conditions under which cell lines are derived and the components that make up both culture media and freezing solutions [193, 194]. Moreover, xeno-free culture media (albeit still containing human serum albumin) have been formulated [195]. A wide range of commercially available cryoprotectant solutions are available [196], including CE marked DMSO, many of which will be suitable for clinical application. Commercial preparations have a number of advantages from a GMP perspective in that they will be sterile and batch tested; though not all will have been certified for other than in vitro use. In the tissue banking area in the UK, regulatory agencies require, at a minimum, that the cryoprotectant is a sterile formulation, and the use of CE marked, sterile DMSO is required.

Commercially available cryoprotectants therefore have the advantage over in-house preparations in that extensive sterility and endotoxin testing will have already been carried out. The use of commercially available freezing and post-thaw washing solutions has been reported for hES and iPS cells [197]. A proprietary solution (STEM-CELLBANKER™) was compared to a standard freezing procedure with 10% DMSO. The commercially available cryoprotectant is a mixture of DMSO, glucose and a high molecular weight polymer (undisclosed) in phosphate-buffered saline. Post-thaw recover was substantially increased without any detrimental impact on proliferation or differentiation, and the freezing protocol could be combined with protocols for derivation and culture in xeno-free conditions, making the cell lines suitable for clinical application.

The fact that at least one component is undisclosed however highlights a potential problem in the use of commercial formulations: that of disclosure. For human application, a risk assessment which includes the components of culture and freezing solutions must be undertaken to assure that adverse events attributable to these solutions are avoided. The use of proprietary solutions in which not all components are declared poses risks. However, the use of confidentiality agreements with the manufacturer permitting disclosure may mitigate this risk.

Scale-Up

Automated culture systems for large-scale production of stem cells are already available [198, 199], and much research is underway to translate laboratory-based cell culture protocols to automated systems; however cryopreservation procedures lag behind. The problems experienced with conventional cryopreservation protocols and the use of straws have undoubtedly contributed to this. Nevertheless novel systems allowing scale-up and cryopreservation have recently been reported in the literature, and some of these are commercially available.

Expansion of hES cells on microcarrier beads in stirred culture systems with conventional cryopreservation of bead-adherent hES cell colonies has been reported [200]. Recovery of hES cells, frozen on feeder cell-seeded microcarriers, was compared with hES cells frozen as freely suspended colonies. A 1.5- to 2-fold increase in recovery was observed, and the cells remained undifferentiated in culture post-thaw.

A commercially available, gas-permeable cassette system for scale-up has also been developed. These ported cassettes are compatible with DMSO (at least in the concentrations used in cryopreservation) and with storage under vapour-phase liquid nitrogen. Since they are gas permeable, the cassettes can be used for the culture of hES cells, and hES cell colonies can be cryopreserved in situ by the simple expedient.
of changing the culture media for the cryoprotectant [201]. In the study using these cassettes, cells were cryopreserved by passive cooling in 10% DMSO and serum. Even under these conditions proliferation ratios, relative to colony fragments cryopreserved in suspension in cryovials, were in the order of 20- to 200-fold greater.

Cryopreservation of mammalian cell lines in cryocyte bags has been reported [202], but the cassette system which allows the manipulation of solutions through scalable ports (and is thus similar in principle to systems used for haematopoietic stem cell preservation) would be a suitable for double bagging and controlled rate cooling.

The use of controlled rate cooling machines within a clean room environment is a potential source of contamination as the nitrogen gas used to cool the samples is vented into the clean room. Cells are thus usually transported out of this environment for freezing. This can lead to potential delays and may contribute to cryoprotectant toxicity if delays are lengthy. Recently, controlled rate cooling devices suitable for use within a cleanroom environment have become available and have been used to freeze embryonic stem cells [203].

Low Temperature Storage and Transportation

The problem associated with low temperature storage and transportation from a GMP perspective is mainly two-fold: maintenance of a suitably low temperature and prevention of contamination.

Storage Temperature

Whilst the therapeutic products of stem cells may not require extended periods of storage, the master and working cell banks from which the products will be derived are likely to require long-term storage. This will require the storage at least below the glass transition temperature required to arrest molecular processes. This has generally been achieved by storage in the liquid or vapour phase of liquid nitrogen. Mechanical refrigeration, delivering stable temperatures below –135 °C, is now reliable, and studies on stem cells indicate no significant differences between this and liquid nitrogen at least in the medium term – up to 5 years [204]. In some systems, temperature fluctuations around the glass transition can be damaging [205], and the storage of vitrified stem cell material at these temperatures, with the possibility of devitrification and ice crystal formation, is probably ill-advised.

Traditionally, storage at ultra-low temperatures was undertaken by immersing the product in liquid nitrogen to ensure as stable a temperature as possible. Following an outbreak of hepatitis in the 1990s in the UK, which was traced to contamination of stem cell harvests during storage under liquid nitrogen [138], this practice was questioned at least for clinically relevant material, and the practice has largely changed to storage in liquid nitrogen vapour (or more accurately gas) phase; a practice also recommended for gametes and embryos [206]. The argument against vapour phase storage, particularly for vitrified material, is that storage vessels designed for under-liquid storage show a considerable vertical temperature gradient when combined with vapour phase platforms. Temperatures as high as –100 °C have been recorded at the top of the inventory system in such vessels. However, a number of technical solutions to this, some designed to be retro-fitted to older storage vessels, have been developed.

The simplest of these is heat shunt. Made either by standing a thermally conducting material such as an aluminium rack system in the liquid present in the base of the vessel [207] or by immersing a copper cooling fin into the liquid [208], the temperature gradient can be dramatically reduced giving temperatures under the lid of the liquid nitrogen storage refrigerator of below –160 °C. Changes to the design of liquid nitrogen storage refrigerators, with the introduction of ‘isothermal’ or ‘dry’ vessels, which have liquid nitrogen-jacketed storage compartments and vessels which are almost completely vacuum insulated have reduced temperature differentials between the bottom and the top of the vessel even more; storage temperature below –180 °C can now be expected even at the top of the inventory system. From a GMP perspective, continuous temperature monitoring and alarming of the storage system is an absolute requirement.

Contamination

The risk of contamination of cell products during low temperature storage are mainly those associated with storage under liquid nitrogen. Vapour phase storage largely overcomes this problem, and experimental studies evaluating the potential for cross-contamination confirmed this [209]. However, the products are still in contact with the external environment and the potential for contamination exists. In haematopoietic stem cell preservation, secondary systems (double bags) are used to protect the primary container. However such systems do not exist currently for non-haematopoietic stem cells. From this point of view, the use of open straws, such as those used to vitrify hES cells, presents problems and is unlikely to be approved by regulatory authorities.

As conventional cryopreservation protocols for slow cooling are improved, the use of cryovials again becomes a realistic possibility. Currently, most cryovials do not prevent the ingress of liquid nitrogen and are therefore potentially open to contamination. Partial sealing of cryovials has been accomplished with heat-sealable membranes [210], and novel closed system cryovials are under development. Recently, the use of pharmaceutical grade plastic cryovials, currently being used to deliver biological therapeutics such as monoclonal antibodies, was evaluated for use in the cryopreservation and storage of cell therapy products including MSCs [211]. The vials were found to be suitable for low temperature storage and transportation of MSCs as well as for use in automated filling systems.
Clinical Application

Cryopreservation of Human Stem Cells for Application

Low Temperature Transportation

For non-vitrified material transportation on pallets of solid CO₂, is the accepted practice. From a regulatory perspective, the containers used for transportation should be validated and the environment within the containers monitored. This may be by individual temperature logging devices or by the use of chemicals which act as thermal exposure indicators, or other devices.

To avoid devitrification, vitrified cells should be transported in liquid nitrogen dry shippers. These come in a range of sizes from those suitable for use internally within a laboratory or cell bank to those for transporting cells over long distances. They function by retaining liquid nitrogen within a molecular sieve-like material which absorbs the liquid nitrogen and permits storage at liquid nitrogen temperatures for up to 14 days. All require charging with liquid nitrogen before use. For compliance with GMP these too need to be validated and periodically re-inspected to ensure that they retain their thermal properties. Data logging lids which monitor and record dry shipper temperature during transportation are available for these containers. Potential contamination and subsequent decontamination are issues for these containers since they are used and re-used many times. Recent studies investigating the potential for contamination have shown that dry shippers do not pose a cross-contamination threat in the absence of damage to the shipment [212] and that if contamination of the shipper does occur they are amenable to decontamination with the method of decontamination dependent on the type of absorbent used to hold the liquid nitrogen [213].

Conclusion

Cryopreservation is a small part of the process of producing stem cells and their derivatives for therapy. While the process is routine for haematopoietic stem cells and largely worked out for MSCs, at least for autologous use, cryopreservation of hES and iPS cells is only now really being explored. The improvement in conventional slow cooling protocols and the novel container systems being developed are likely to provide systems that are compatible with the requirements of GMP, regulation, automation, and scale-up. However, the research being undertaken is still empirically based, and sound methodological approaches to optimising cryobiological variables are still required if the maximum benefits for regenerative medicine therapies are to be realised.

Disclosure Statement

The author declared no conflict of interest.

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Cryopreservation of Human Stem Cells for Clinical Application


