Cryomicroscopic Investigations of Freezing Processes in Cell Suspensions
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Abstract: This study evaluates the freezing response of three different cell types, Pacific oyster embryos (~50 μm in diameter), Jurkat cells and HeLa cells (~12 to 15 μm’s in diameter), using cryomicroscopy. Freezing experiments were performed on oyster embryos at cooling rates of either 5 or 10 °C/min, while Jurkats were cooled at either 1 or 10 °C/min and HeLa cells were cooled at either 1, 15 or 20 °C/min, respectively. The experiments with oyster embryos were primarily designed to investigate the phenomena of intracellular ice formation (IIF) while the experiments for Jurkat and HeLa cells were designed to investigate both cellular dehydration and IIF during freezing. IIF was characterized by the abrupt black flashing during the cooling steps while the cellular dehydration experiments were characterized by the volumetric (projected area) shrinkage of the cells during the cooling steps. Mathematical models were fit to the cellular dehydration data to obtain the Jurkat and HeLa cell membrane permeability ($L_{pm}$) at the reference temperature (273.15 K), the apparent activation energy of the cellular dehydration process ($E_{ac}$) or the temperature dependence of $L_{pm}$. The temperature dependence of IIF in the oyster embryos, the Jurkat cells and the HeLa cells were also determined.

Keywords: Intracellular ice formation, water transport or cellular dehydration, pacific oyster embryos, HeLa cells, jurkat cells.

INTRODUCTION

Cryopreservation refers to the technique of storing a living organism at extremely low temperatures in suspended animation, for long periods of time, so that it may be revived and restored to the same living state. Cryopreservation techniques have already been shown to be useful in a variety of mammalian systems such as lymphocytes, platelets, granulocytes, gametes, hepatocytes, bone marrow stem cells, cornea and skin, pancreatic tissue, human and non-human mammalian oocytes, tissue engineered equivalents, adult stem cells, insects, fish, micro-organisms and algae, rat and human liver slices [1-12]. Alternatively, as demonstrated by Dr. James Arnott and co-investigators in the 1850s, cold temperatures could also be used for the treatment of malignant tumors, a process termed as cryosurgery [13-20].

Both cryopreservation and cryosurgery require an understanding of the relationship between the biological system being frozen and the applied cooling stress. When the cell in a solution is subjected to a cooling protocol, the initial ice formation takes place in the extracellular space. The cell membrane acts as a barrier and prevents ice formation inside the cell resulting in super-cooling of the intracellular water. The unfrozen super-cooled intracellular water remains at a temperature higher than the partially frozen extracellular solution [20-24]. This leads to thermodynamic nonequilibrium and provides the driving force for the two biophysical processes during freezing, either cellular dehydration (water transport) or intracellular ice formation, IIF [22-26]. Extensive data suggests that at “slow” cooling rates water transport is the dominant mechanism while IIF is the dominant mechanism at “rapid” cooling rates [25-30]. Both cellular water transport and IIF have been proved to be deleterious to the post-thaw survival of biological systems and an optimal cooling rate exists between these two extremes [24-32]. For example, when the cooling rate is slow, the intracellular water has enough time to flow out of the cell through the semi-permeable membrane and join the extracellular space. This results in changes in the extracellular region, such as in pH and ionic concentrations and potentially causes the tertiary structure of proteins to unfold. Hence, some of the original properties of the cell (and its internal structures) are either diminished or eliminated [20-32]. Additional damage may also take place through the mechanical interactions between the extracellular ice crystals and cells [33]. On the other hand, IIF is generally considered to be lethal as it causes injury to the intracellular structures and the cell membrane [20-32]. Thus, the balance between the membrane’s water permeability and the probability of IIF results in different optimal cooling rates for different cells, i.e. the exact magnitude of the “slow” and “rapid” cooling is dependent upon the cell type and needs to be experimentally determined.

In this work, the responses of three different cell types (Pacific oyster embryos, Jurkat and HeLa cells) to an imposed cooling stress were evaluated using cryomicroscopy. Prior cryopreservation studies on aquatic cells (gametes and embryos) are quite extensive and were conducted to assist selective breeding programs [34-41]. For example, Salinas-Flores et al. [42], determined the membrane permeability characteristics of Pacific oyster oocytes and the development of optimized methods to add and remove a cryoprotective agent (ethylene glycol). Lin et al. [43] studied the effects of cooling rate, choice of cryoprotectant, and the seeding temperature on the survival of oyster embryos. The second cell type used in this study the Jurkat cells are routinely used to study T cell leukemia, T cell signaling and the expression of...
various chemokine receptors [44]. But prior cryobiological work on Jurkat cells is limited [45]. There is an increasing need to understand the low temperature behavior of Jurkat cells not only for immunological reasons but also for the application in adoptive tumor therapy [46]. And finally, like Jurkat cells, HeLa cells are also an immortal cell line derived from cervical cancer cells [47]. Similar to Jurkat cells, these cells were also chosen for analysis because of their robustness, easy availability, and because they are relatively less understood among the cancer cells.

**MATERIALS AND METHODS**

**Cryomicroscopy Experiments**

Cryomicroscopy involves application of cryogenic temperatures to cellular systems mounted under a light microscope to study the biophysical response of cells to freezing. This technique has been refined and modified over years to study the dynamic response of a variety of plant and mammalian cells under various freezing conditions [48-62]. Some of the important advances in cryomicroscopy include the mounting of a cold stage on a microscope instead of cooling the entire microscope; the use of a digital control system in lieu of the less accurate analog control system; the use of video recordings to record the events during freezing; and the use of computer enhanced image analysis techniques to overcome the low resolution of the video recordings. Thus, the currently available cryomicroscopy systems integrate the functions of sample temperature control, image storage & event correlation, and image analysis. Cryomicroscopy can, therefore, be used to obtain images of a dehydrating cell during the course of the freezing protocol. Hence, by calculating the change in surface area of a cell from its projected two dimensional image one can extrapolate it to the three dimensional change in cell volume (cellular dehydration) as a function of cooling rate and sub-zero temperature [48-62]. Cryomicroscopy also helps in characterizing IIF through visual inspection of the blackening of cells due to scattering of light by opaque ice inside the nucleated cells [48-62].

The cryomicroscopy experimental procedures utilized in the present study are similar to those described extensively in the literature [48-62] and were performed using a commercially available BCS-196 cryostage (Linkam Scientific™, Surrey, UK). The Linkam™ cryostage is capable of controlled cooling and heating between –125 ˚C and +160 ˚C at cooling/heating rates ranging from 0.1 to 130 ˚C/min (Fig. 1A). Typically, the cryostage is cooled by a constant flow of liquid nitrogen; thus, equilibrating the cryostage at ~ –160 ˚C. The cryostage temperatures are measured by platinum resistance thermocouples. A temperature controller maintains the cryostage at the desired pre-set temperature by utilizing electrical (power) heating, i.e., the amount of cooling due to the liquid nitrogen flow is adjusted and counterbalanced with electrical heaters. Hence, maintaining the cryostage at the desired temperature or set-point. A time varying temperature (or a desired cooling rate) can be created using temperature-control software that dynamically changed the amount of heating, i.e., the sample could either be cooled or heated by either decreasing or increasing the amount of heating. The cryostage (Fig. 1B) was mounted with a circular quartz crucible to hold the cell suspension during freezing. A cover-slip was placed on top of the cell suspension within the quartz crucible to prevent leakage/media evaporation and to ensure proper visibility of the cells during the freezing process. The Linkam™ cryostage was attached to a microscope (Nikon Eclipse E600, Nikon Instruments, NY) fitted with a Photometrics Coolsnap cf camera (Hamamatsu, Photonics, Bridgewater, NJ). During the experiments a live video signal was sent from the camera to the attached DELL personal computer and the images recorded with the help of commercially available Metamac software (Universal Imaging Corp., Buckinghamshire, UK).

**Image Analysis**

Image analysis of cells involved the evaluation of volumetric shrinkage of cells due to dehydration and evaluation of number of cells that underwent intracellular ice formation. Evaluation was based on the assumption that the cells were regular and spherical. The change in the diameter of the cell was measured after a dehydration experiment which was then utilized in evaluating the change in corresponding cellular volume [48-62]. For IIF experiments, the images obtained through the camera were stacked using the Metamac software to make movies. Usually a cluster of around 30-40 cells were focused under the microscope and were examined for blackening. The number of cells that turned black due to IIF at different temperatures through the protocol was noted. Percentage intracellular ice formation (P-IIF) was evaluated as the ratio of the number of cells that turned black due to IIF to the total number of cells and was plotted against temperature for different freezing rates.

**Pacific Oyster (Crassostrea gigas) Embryos**

Oyster embryos were obtained from a population of fertilized eggs, about 120 minutes after the artificial fertilization in sea water of 34+2% salinity at a temperature of 27±1 ˚C. Hundreds of oyster embryos were pooled from three females. A 10x D-PBS (Dulbecco’s Phosphate Buffer Saline) solution was used as the cellular suspension media. The embryos were placed on a petri-dish to be examined under the cryomicroscope for an applied cooling stress. A group of embryos were focused under the cryomicroscope, and a predetermined freeze thaw protocol was used to examine their freezing response. The main aim of the experiment was to identify the freezing rates at which the cells showed darkening or flashing to reveal intracellular ice formation. The freezing protocol used for the oyster embryo experiments was:

**Step 1:** Cool the embryos from 23 ˚C to 4 ˚C at 5 ˚C/min

**Step 2:** Cool the embryos from 4 ˚C to ~ –6 ˚C at 1 ˚C/min. Note that the phase change temperature of 10x D-PBS is ~ –5.6 ˚C.

**Step 3:** Cool the embryos from ~6 ˚C to ~40 ˚C at either 5 or 10 ˚C/min.

**Step 4:** Heat the embryos from ~40 ˚C to 20 ˚C at either 5 or 10 ˚C/min.

**Jurkat Cells**

Jurkat cells were maintained in 25 cm² T-flasks (BD Falcon, Franklin Lanes, NJ) with 5 ml of HyQ® RPMI-1640-reduced serum medium (RPMI-RS) supplemented with 3% fetal bovine serum (FBS) and incubated at 37 ˚C in a humidified
atmosphere containing 5% CO₂. For cryomicroscopy experiments, Jurkat cells were transferred to a centrifuge tube and were spun down at 1900 rpm for 5 minutes using Eppendorf Minispin Plus centrifuge. The pellet was resuspended in 1X Dulbecco’s Phosphate Buffered Saline (DPBS) solution, which was considered isotonic for these cells. Experiments on Jurkat cells were performed at either 1 or 10 °C/min in order to observe their response, at both “slow” and “fast” freezing rates. A typical freezing protocol used for the Jurkat cell experiments was:

**Step 1:** Cool the Jurkats from 23 °C to 4 °C at 10 °C/min

**Step 2:** Cool the Jurkats from 4 °C to −1 °C at 1 °C/min and equilibrate at −1 °C for 1 min to facilitate extracellular ice nucleation. Note that the phase change temperature of 1x D-PBS is ~ −0.6°C.

**Step 3:** Cool the Jurkats from −1 °C to −30 °C at either 1 or 10 °C/min.

**Step 4:** Heat the Jurkats from −30 °C to 20 °C at 10 °C/min.

**HeLa Cells**

HeLa cells were maintained in 25 cm² flasks (BD Falcon, Franklin Lakes, NJ) with 5 ml of Dulbecco’s Modified Eagle’s medium-reduced serum (DMEM-RS) supplemented with 3% fetal bovine serum (FBS) and incubated at 37 °C in a humidified atmosphere containing 5% CO₂. HeLa cells were plated at a density of 80,000 cells per cm² in 12 well cultured plates (BD Falcon, Franklin Lakes, NJ) and were allowed to adhere and grow for 24 h prior to trypsinization (0.25% Trypsin). In order to prepare the cells for experiments, the following steps were followed: i) aspiration of cell growth media from T-flask to waste; ii) addition of 5 ml of 1x D-PBS; iii) addition of 5 ml of Trypsin and equilibration for 10 mins. This facilitates the detachment of the HeLa cells from the T-flask substrate; iv) addition of 5 ml of DMEM-RS (reduced serum) with 3% Fetal Bovine Serum (FBS) to prevent further deleterious trypsinization; v) transfer of HeLa cells suspension to a 50 ml centrifuge tube; vi) spin down of cells at 2400 rpm for 5 min using an Eppendorf Minispin Plus centrifuge; vii) decant off the supernatant and...
re-suspend the HeLa cells in 1x D-PBS. Experiments on HeLa cells were performed at either 1, 15 or 20 °C/min in order to observe their response, at both “slow” and “fast” freezing rates. A typical freezing protocol used for the HeLa cell experiments was:

**Step 1:** Cool the HeLas from 23 °C to 4 °C at 10 °C/min

**Step 2:** Cool the HeLas from 4 °C to –1 °C at 1 °C/min and equilibrate at –1 °C for 1 min to facilitate extracellular ice nucleation. Note that the phase change temperature of 1x D-PBS is ~0.6 °C.

**Step 3:** Cool the HeLas from –1 °C to –30 °C at either 1, 15 or 20 °C/min.

**Step 4:** Heat the HeLas from –30 °C to 20 °C at 10 °C/min.

**Water Transport Model for Jurkat and HeLa Cells**

The general model of water transport during freezing in single cell systems is reviewed elsewhere [63]. The model predicts the change of the cell volume with decreasing temperature T after ice has formed outside of an unfrozen cell. This is depicted mathematically as:

\[ dV/dT = L_p ART (C_i - C_o) / B, \]

where \( L_p \) is the water permeability of the cell plasma membrane, \( A \) is the surface area of the cell, \( R \) is the gas constant, \( C_i \) and \( C_o \) are the concentrations (osmotic potentials) of the intracellular and extracellular (unfrozen fractions) respectively, and \( B \) is the cooling rate of the protocol (°C/min). The cell membrane permeability, \( L_p = fn(L_{pg}, E_{lp}) \), itself depends on two biophysical parameters, \( L_{pg} \), the reference water permeability at the phase change temperature of water, and \( E_{lp} \), the activation energy for the water transport process [64].

Since the temperature, time, geometry and concentrations are all known in the experiment, the only unknowns in the equation are \( L_{pg} \) and \( E_{lp} \) which are called the biophysical parameters of water transport. These parameters (\( L_{pg} \) and \( E_{lp} \)) can be determined by a curve fit of the water transport equation to the experimental data as previously described [9-11, 30-32, 65, 66]. Once determined, these biophysical parameters can be incorporated into a recently developed Generic Optimal Cooling Rate Equation (GOCRE) that defines the optimal cooling rate \( B_{opt} \) (in °C/min) as [66]:

\[ B_{opt} = 1009.5 \exp ^{-0.0564E_{lp} } \cdot (L_{pg}) \cdot (\frac{A}{WV}). \]

In the GOCRE, \( L_{pg} \) and \( E_{lp} \) represent the membrane permeability parameters (in μm/min-atm and Kcal/mole, respectively), while the term \( A/WV \) (in μm⁻¹) represents the ratio of the available surface area for water transport (A) to the initial volume of intracellular water (WV = \( V_o - V_h \); \( V_o \) is the isotonic or initial cell volume and \( V_h \) is the osmotically inactive cell volume).

**RESULTS AND DISCUSSION**

**Pacific Oyster Embryos (IIF Measurements)**

During the cryomicroscopy experiments the embryos were checked for reduction in size or diameter which would have indicated dehydration or water transport. In addition, the embryos were also checked for IIF through sudden blackening or darkening. Cells undergoing IIF usually dem-onstrate darkening because of ice formation that deflects or restricts the passage of light through them. Thus, transforming them from transparent to opaque cells. During a typical experiment (See Fig. 2), a total of 40 oyster embryos were studied. The oyster embryos were found to be spheres with a diameter of ~60 μm. Fig. (2A) shows an image that was taken at the beginning of the experiment and at room temperature (~23°C). The oyster embryos are clearly visible and appeared to be semi-transparent and the extracellular solution of 10x D-PBS also appear as a clear and transparent fluid. Fig. (2B) shows an image of the same cells (as in Fig. 2A) but now cooled to ~15 °C at 5 °C/min. Notice that the extracellular solution is now frozen and that some of the oysters are exhibiting IIF or darkening. Figs. (2C and 2D) represent the same cell population cooled at 5 °C/min to ~18 °C and ~19.9 °C, respectively. Notice the progressive increase in the percentage of embryo’s exhibiting IIF, i.e. cells that are exhibiting “blackening” or “darkening”. Percentage of embryos exhibiting IIF (P-IIF) was evaluated as the ratio of the number of embryos that darkened (exhibited IIF) to the total number of embryos that were present in the field of view (See Fig. 3). As shown in Fig. (3), oyster embryos cooled at 5 °C/min underwent IIF between ~6 °C and ~20 °C while embryos cooled at 10 °C/min exhibited IIF between ~6 °C and ~18 °C. Note that the cumulative fraction (%) of cells exhibiting IIF was obtained from 8 to 10 separate experiments for each cooling rate with approximately 10 cells in the field of view per experiment and pooled as a single population per cooling rate. Since, the data is a representation of the cumulative fraction of cells experiencing IIF from all the experiments (or from a population of cells). Hence, no error bars are shown within Fig. (3).

It is, generally believed that IIF is a deleterious event, that is lethal to cells. Hence, knowledge of freezing rates that induce IIF in oyster embryos will reveal the range of “fast” cooling rates; in our studies these are cooling rates ≥ 5 °C/min. Thus, in the absence of any cryoprotective agents it is essential that oyster embryos be cooled at rates slower than 5 °C/min, possibly as low as 1 or 2 °C/min. Unfortunately, additional experiments could not be conducted at the “slower” rates due to the restricted and seasonal availability of oyster embryos.

**Relevance to Oyster Embryo Cryopreservation**

Recent United States Department of Agriculture (USDA) statistics indicate that the commercial value of oyster embryos produced within the continental United States is greater than $100 million per year. However, the total production has decreased from 1998 due to diseases and adverse weather conditions while, the consumption of oysters has jumped from an average of 22 pounds per person to ~31 pounds per person in the US. Thus, over the last decade, many researchers have made attempts to improve oyster hatcheries. Obviously newer methods of hatchery production needed to be developed in order to keep up with the growing demand. One possible solution might lie in developing successful cryopreservation protocols for preserving oyster spermatozoa and oocytes/embryos and enabling fertilization. Although there were some exceptions [34-43,67-69] the study of cryobiology was largely neglected in most aquatic species. For example, Salinas-Flores et al. [68] determined that the Pacific oyster oocytes behaved as ideal osmometers...
Fig. (2). Photomicrographs of Pacific oyster embryos being cooled at 5 °C/min are shown at various temperatures. Specifically, at room temperature or 23 °C (2A), –15 °C (2B), –18 °C (2C) and –19.9 °C (2D) are shown. Note the progressive increase in the number of cells exhibiting blackening or darkening associated with intracellular ice formation (IIF) as the sub-zero temperature is reduced, as shown in Figs. (2B to 2D). The formation of extracellular ice is evident in Figs. (2B to 2D).

Fig. (3). The relationship between the probability of intracellular ice formation (P-IIF) and sub-zero temperature is shown for Pacific oyster embryos at two different cooling rates (5 and 10 °C/min). The unfilled columns represent the data at 5 °C/min while the shaded columns represent the corresponding data at 10 °C/min. The % of oyster embryos exhibiting IIF is shown on the y-axis while the sub-zero temperature (°C) is shown on the x-axis. Note that the cumulative fraction (%) of cells exhibiting IIF is presented as a population based measurement and the data is a representation of the cumulative fraction of cells experiencing IIF from all the experiments (or from a population of cells). Hence, no error bars are shown within Fig. (3).
with an inactive cell volume ($V_b$) of 0.48$V_o$ while Tervit et al. [69] reported that ethylene glycol (10%) and dimethyl sulfoxide (15%) were the most effective cryoprotectants for oyster oocytes resulting in high post-thaw fertilization rates of ~51% and ~45%, respectively. Tervit et al. [69] also suggested that the use of Milli-Q water instead of sea water as a base medium, and a 5 minute isothermal hold at either −10 °C or −12 °C, significantly improved the Oyster oocyte fertilization rates. The data presented in this study suggests that Pacific oysters embryos need to be cooled at rates of ~1 to 2 °C/min for optimal post-freeze response. This value is comparable to those published in the literature for mouse and human embryos [70].

**Jurkat Cells (Water Transport Measurements)**

Assuming a Jurkat cell to be spherical, the initial (pre-frozen) diameter of a Jurkat cell was noted as 12 ± 1.1 μm [45]. To account for any possible variations in the cell diameter between Jurkat cells, the normalized volume of a given Jurkat cell at a given sub-zero temperature was evaluated as a ratio of the experimentally determined diameter at this (sub-zero) temperature to the original diameter of the same cell at the initial (pre-frozen) temperature of 23 °C. Any reduction in diameter can be attributed to corresponding reduction in the Jurkat cell volume or to the process of cellular dehydration (water transport). For example, Figs. (4A, 4B and 4C) represent a Jurkat cell being cooled at 1°C/min at ~23°C, −2 °C and −7 °C, respectively. The corresponding reduction in volume as a function of sub-zero temperature was analyzed and plotted, as shown in Fig. (4D). As shown in Fig. (4D), the cellular dehydration process in Jurkat cells terminates at approximately −7 °C and the final (fully) dehydrated volume or the osmotically inactive cell volume ($V_b$) was 0.52$V_o$. This final osmotically inactive volume for Jurkat cells is in good agreement with earlier published results [45]. By curve-fitting the data shown in Fig. (4D) to the water transport model described earlier, we determined that the $L_{pg}$ of Jurkat cells is 0.20 μm/min-atm while the $E_{lp}$ is 69.7 Kcal/mole (the goodness of fit was 0.96; note that a value of 1 indicates a perfect fit between the experimental data and the numerical simulation). Utilizing the GOCRE, these values translated into an optimal cooling rate of ~ 2.2 °C/min for Jurkat cells.

![Fig. (4A)](image1.png) ![Fig. (4B)](image2.png) ![Fig. (4C)](image3.png) ![Fig. (4D)](image4.png)

**Fig. (4).** Photomicrographs of Jurkat cells being cooled at 1 °C/min are shown at various temperatures. Specifically, at room temperature or 23 °C (4A), −2 °C (4B) and −7 °C (4C) are shown. Note the progressive decrease in the cell size as the subzero temperature is decreased and the associated loss of intracellular water. Fig. (4D) shows the relationship between the non-dimensional cell volume and the sub-zero temperature for Jurkats cooled at 1 °C/min. The filled circles represent the experimentally determined non-dimensional diameter at this (sub-zero) temperature to the original diameter of the same cell at the initial (pre-frozen) temperature of 23 °C. Any reduction in diameter can be attributed to corresponding reduction in the Jurkat cell volume or to the process of cellular dehydration (water transport). For example, Figs. (4A, 4B and 4C) represent a Jurkat cell being cooled at 1°C/min at ~23°C, −2 °C and −7 °C, respectively. The corresponding reduction in volume as a function of sub-zero temperature was analyzed and plotted, as shown in Fig. (4D). As shown in Fig. (4D), the cellular dehydration process in Jurkat cells terminates at approximately −7 °C and the final (fully) dehydrated volume or the osmotically inactive cell volume ($V_b$) was 0.52$V_o$. This final osmotically inactive volume for Jurkat cells is in good agreement with earlier published results [45]. By curve-fitting the data shown in Fig. (4D) to the water transport model described earlier, we determined that the $L_{pg}$ of Jurkat cells is 0.20 μm/min-atm while the $E_{lp}$ is 69.7 Kcal/mole (the goodness of fit was 0.96; note that a value of 1 indicates a perfect fit between the experimental data and the numerical simulation). Utilizing the GOCRE, these values translated into an optimal cooling rate of ~ 2.2 °C/min for Jurkat cells.
Jurkat Cells (IIF Measurements)

As shown in Fig. (5A), Jurkat cells cooled at 10 °C/min to −18 °C exhibited a P-IIF of 100%, i.e., all the Jurkat cells experienced IIF when cooled at 10 °C/min. Jurkat cells experienced IIF within the narrow temperature band between −16 and −18 °C, when cooled at 10 °C/min. A plot of P-IIF with sub-zero temperature for Jurkats cooled at 10 °C/min is shown in Fig. (5B). Note that the cumulative fraction (%) of cells exhibiting IIF is presented as a population based measurement and the data is a representation of the cumulative fraction of cells experiencing IIF from all the experiments (or from a population of cells). Hence, no error bars are shown within Fig. (5B).

HeLa Cells (Water Transport Measurements)

Assuming a HeLa cell to be spherical, the initial (pre-frozen) diameter of a HeLa cell was noted as 15 ± 1.5 μm [45]. The water transport analysis for HeLa cells was performed as described earlier for Jurkat cells at a cooling rate of 1 °C/min. The corresponding reduction in HeLa cell volume as a function of sub-zero temperature was analyzed and plotted, and is shown in Fig. (6A). As shown the cellular dehydration process terminates at approximately −7 °C and the final (fully) dehydrated volume was 0.45V₀. This final osmotically inactive volume for HeLa cells is in good agreement with earlier published results [45]. By curve-fitting the data shown in Fig. (6A) to the water transport model described earlier, we determined that the Iₚₚ of HeLa cells is 0.05 μm/min-atm while the Eᵦ is 20.9 Kcal/mole (the goodness of fit was 0.97). Utilizing the GOCRE, these values translated into an optimal cooling rate of ~6.5 °C/min for HeLa cells.

HeLa Cells (IIF Measurements)

A plot of P-IIF with sub-zero temperature for HeLas cooled at 15 and 20 °C/min is shown in Fig. (6B). As shown in Fig. (6B), HeLa cells cooled at 15 °C/min exhibited IIF between −2 and −11.5 °C while the HeLa cells cooled at 20 °C/min exhibited IIF between −2 and −19.2 °C, respectively. Note that all the HeLa cells did not experience IIF when cooled at either 15 or 20 °C/min. Approximately, ~81% and ~94% of the HeLa cells exhibited IIF when cooled at 15 and 20 °C/min, respectively. Additional experiments at 10 °C/min with HeLa cells did not show any evidence of IIF (data not shown).

Relevance of Jurkat and HeLa Cell Measurements to Cryosurgery

The results presented here are qualitatively similar to those of other cells using cryomicroscopy [48-62] and more specifically to earlier studies on HeLas [50]. Briefly, at low cooling rates, cellular dehydration occurs and as the cooling rate is increased the percentage (and the probability) of cells exhibiting IIF increases. The importance of the P-IIF to cryobiology is related by an observation by Toner [70] that the cooling rate that induces 50% of the cells to exhibit IIF is optimal, i.e., cooling rates that are above or below this cooling rate are sub-optimal for cryopreservation but conversely, are optimal for cryodestruction. Hence, the P-IIF data obtained for Jurkat cells in the present study indicate that the extensive cellular damage due to IIF will occur when these are cooled > 5 °C/min while the corresponding cooling rates for HeLa cells are > 10 °C/min. Conversely, the water transport data indicates that both these cell types need to cooled at < 1 °C/min for extensive solute effects or slow cooling injury to occur. These damaging cooling rates (< 1 °C/min and > 5 °C/min for Jurkats; < 1 °C/min and > 10 °C/min for HeLas) are present within a typical cryosurgical ice-ball, i.e., at regions at the edge of the frozen region or the ice-ball (cooling rates < 1 °C/min) and at regions close to the inserted ice probe (cooling rates > 5 °C/min). Obviously, the behavior and response of these cells when they are embedded in an extra-cellular tissue matrix (like in a tumor tissue) will be dissimilar to isolated cells, i.e. the cell types used in the present study [71]. And finally, the membrane and protein denaturation during freezing/thawing and the associated injury...
might also play a role in determining the outcome of a cryo-
surgical protocol [72].

CONCLUSIONS

Freezing experiments were performed on three different
types of cells, Pacific oyster embryos, Jurkat cells and HeLa
cells. In this work, IIF through darkening of the cells was
observed in Pacific oyster embryos cooled at 5 and 10
˚C/min. The water transport model-simulated cooling response at 1
˚C/min is also shown as a solid line in Fig. (6A). In Fig. (6A), the non-dimensional HeLa cell volume is shown on the y-axis while the sub-zero temperature
(˚C) is shown on the x-axis. Fig. (6B) shows the relationship between the probability of intracellular ice formation (P-IIF) and sub-zero tem-
perature (˚C) for HeLa cells being cooled at either 15 or 20 ˚C/min. In Fig. (6B), the unfilled columns represent the data at 15 ˚C/min while the shaded columns represent the corresponding data at 20 ˚C/min. In Fig. (6B), the % of HeLa cells exhibiting IIF is shown on the y-axis while the sub-zero temperature (˚C) is shown on the x-axis. Note that the cumulative fraction (%) of cells exhibiting IIF is presented as a population based measurement and the data is a representation of the cumulative fraction of cells experiencing IIF from all the experiments (or from a population of cells). Hence, no error bars are shown within Fig. (6B).

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Cryomicroscopy of Cell Suspensions


