INTRODUCTION. Vitrification is the solidification of a liquid brought about not by crystallization but by an extreme elevation in viscosity during cooling [1]. It is a widely applied alternative to standard slow programmable freezing methods for cryopreservation because of the higher survival rates of cells after thawing [2, 3]. Vitrification was first proposed in 1985 by Greg Fahy and William F. Rall [4] as a method for cryopreserving complex tissues such as whole organs. The motivation for vitrification was that conventional freeze preservation invariably destroyed organs by disrupting sensitive tissue structures with ice crystals [5]. Adding a high concentration of cryoprotectants can limit the amount of ice that forms during freezing so that less destroying the normal structure of the tissue occurs. But freezing with high cryoprotectant concentrations still causes serious structural damage. Combination of structural damage and toxicity makes recovery of frozen neural tissue impossible with current technology [5].

Vitrification of water requires an ultra-rapid cooling (e.g. 10 million degrees per second for pure water), but can be achieved without cryoprotectants. Only in 2005 the cooling rate required to form a glassy state in pure water was achieved [6]. The necessary cooling rates for solutions were significantly less than for pure liquids. For aqueous solutions of typical cryoprotectants, cooling rates of about 0.1–10 °C/sec (roughly 10–1,000 °C/min) are sufficient to achieve vitrification [7].

Cryoprotectants, or cryoprotective agents (CPAs) have been extensively used for vitrification [16, 17]. CPAs are chemicals which prevent cell damage caused by cryopreservation [8]. Such substances as alcohols, amides, oxides and polymers with corresponding functional groups can be effective CPAs. These chemicals increase the viscosity of aqueous solutions, reduce the freezing point and lower the ice nucleation temperatures of aqueous solutions. Some CPAs can reduce the glass transition temperature of a solution. CPAs from the point of view of ability to cell penetration can be good penetrating (dimethyl sulfoxide, glycerol, propylene glycol, ethylene glycol, methanol), non-penetrating (mono-, di- and polysaccharides; such polymers as polyethylene glycol, polyvinyl pyrrolidone and polyvinyl alcohol) and mixed-action CPAs (polyethylene oxide). For polymeric CPAs there are restrictions on molecular mass due to data about its cryoprotective properties, rheology and toxicity: 100–1500 Da for polyethylene oxide, 12.5–25 kDa for polyvinyl pyrrolidone and 250–500 kDa for hydroxyethyl starch [9].

So, effective CPAs must be low- or non-toxic, well-soluble in water and aqueous solutions. It has to lower the ice nucleation and completely suspend ice crystallization from CPA-water eutectic system. At the same time it should not precipitate salts and proteins from aqueous solutions before transition to vitrified state.

into amorphous state. Unfortunately, some substances (for example, methanol and ethanol) with good physicochemical properties can not be used as CPAs due to high toxicity even in low concentrations.

Main properties of CPAs caused by the presence in their molecules groups that can form hydrogen bounds. Substances that can form a network of hydrogen bonds, such as polyhydric alcohols, significantly increase the viscosity of aqueous solutions, which allow reducing the required concentration of CPA. With the further water freezing CPA solutions became hypertonic, that leads to partial dehydration of cell. In this state intracellular liquid cell media going super cooled and suppress the ice formation.

Usually cryopreservation solutions (mixtures of CPA with some additions) are less toxic than pure CPA. In most cases cryopreservation solutions includes penetrating CPA, non-penetrating CPA (optional ingredient), carrier solution and ice blocker (optional ingredient) [17].

One of the main requirements for penetrating CPA is the ability to rapidly go through the cell membrane and easily removed from it. It reduces osmotic effects of cooling and further warming. Fast penetration of cell by CPA prevents the formation of damaging concentration gradients between extra- and intracellular environment on cell membrane in selected temperature range. Main disadvantage of penetrating CPA is necessity of removing CPA from cells after thawing. In another case the heavy damage of cell membranes and intercellular structures can be derived. CPA ability to penetrate cells depends on molecular mass, temperature, type of cells and properties of cell membranes.

Protective effect of non-penetrating CPA caused by the ability to partially dehydration of cells, that reduces the probability of intracellular ice crystallization. Non-penetrating CPA are usually less toxic than penetrating CPA at the same concentration.

Carrier solution contains salts, osmotic agents, pH buffers, and sometimes nutritive ingredients or apoptosis inhibitors. The ingredients are usually present at near isotonic concentration (300 millimoles) so that cells neither shrink nor swell when held in carrier solution [10].

Ice blockers are synthetic ice interface dopants specifically designed to bind to the basal plane and prism faces of ice crystals (and ice nucleates) [11]. Molecules of ice blockers must have the spacing polar groups at intervals corresponding to the lattice spacing of water molecules on the crystal faces of ice. Low molecular weight polyvinyl alcohol and polyglycerol, called X-1000 and Z-1000, and biological antifreeze proteins are examples of ice blockers [11, 12].

The aim of this study was to find the optimal ratio of penetrating (DMSO) and non-penetrating (saccharose and polyethylene glycols) cryoprotectants to achieve the greatest survival of human somatic cells in the two-step process of vitrification.

METHODS OF RESEARCH. Breast adenocarcinoma cell line MCF-7 was kindly presented by of Bank of cell lines of man and animals R. Ye. Katvetskyi Institute of Experimental Pathology, Oncology and Radiobiology NAS Ukraine. Cells were incubated under standard conditions in 5 % of CO₂, 95 % humidity in RPMI – 1640 medium (Sigma) which was supplemented with 10 % fetal bovine serum (FBS, Sigma, USA) and 40 mg/ml gentamycin (Sigma) for cell culture. Also buffered solution of 1 M HEPES (Sigma) pH 7.4 was used. For washing cells phosphate – saline buffered solution (PBS): 1.5 mM KH₂PO₄, 8.1 mM Na₂HPO₄, 2.7 mM KCl, 140 mM NaCl was used.

Vitrification solutions (VS) on the first stage of investigation were based on saline (0.9 % NaCl) with supplements according to Table 1. VS were used for determination of cooling rate by time-dependent thermometry. Glucose (Mr=180.16 Da), saccharose (Mr=342.30 Da), glycerol (Mr=78.13 Da) and dimethyl sulfoxide (Mr=92.09 Da) were used as cryoprotective agents (Table 1) for estimation of cooling rate at the first stage of our investigation.

Table 1 – Cryopreservation solutions studied on the cooling rate stage

<table>
<thead>
<tr>
<th>Name</th>
<th>Glucose, M</th>
<th>Saccharose, M</th>
<th>Glycerol, %</th>
<th>DMSO, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>VS1.1</td>
<td>0.2</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>VS1.2</td>
<td>–</td>
<td>0.2</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>VS1.3</td>
<td>–</td>
<td>–</td>
<td>5</td>
<td>–</td>
</tr>
<tr>
<td>VS1.4</td>
<td>–</td>
<td>–</td>
<td>2.5</td>
<td>–</td>
</tr>
<tr>
<td>VS1.5</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>5</td>
</tr>
<tr>
<td>VS1.6</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>2.5</td>
</tr>
<tr>
<td>VS1.7</td>
<td>–</td>
<td>0.2</td>
<td>5</td>
<td>–</td>
</tr>
<tr>
<td>VS1.8</td>
<td>–</td>
<td>0.2</td>
<td>2.5</td>
<td>2.5</td>
</tr>
<tr>
<td>VS1.9</td>
<td>0.2</td>
<td>–</td>
<td>2.5</td>
<td>2.5</td>
</tr>
<tr>
<td>VS1.10</td>
<td>0.2</td>
<td>–</td>
<td>5</td>
<td>–</td>
</tr>
</tbody>
</table>
For detection of cell viability after vitrification RPMI (Sigma, USA) medium with 10 % of FBS (Sigma, USA) we used. End concentrations and supplements are demonstrated in Table 2.

**Determination of cooling rate by time-dependent thermometry.**

Boiling liquid nitrogen was cooled to -210 °C by pumping out air and reducing pressure in the Dewar flask. Then the air carefully returned to the Dewar flask and then test-tube with sample (Table 1) and thermo-sensor immersed in liquid nitrogen. Changes of temperature were recorded during 118 seconds, after that test-tube was removed from liquid nitrogen and prepared for the next sample.

**Cell culture studies. Monolayer culture (2-D culture). Study design.**

MCF-7 cells in 2-D culture was cultured five days in full culture medium. For freezing several groups were formed with different condition of vitrification solutions (Table 2). After thawing cell viability was investigated by MTT-assay, light microscopy and tripan blue staining. Micro-photos of 2-D cell cultures were made after 7 days of incubation.

**Analysis of cell viability by MTT – colorimetric assay.**

MTT test was used in order to analyze the impact of vitrification conditions on cell viability. The MTT-test was based on conversion of tetrazolium salts to formasan crystals by NAD(P)H-dependent mitochondrial oxidoreductase enzymes in alive cells. Protocol was described by T. Mosmann [18].

In brief, MCF-7 cells after thawing were incubated in full culture medium during 24 hours. Then it was analyzed with 3-[4,5-dimethylthiazol-2]-2,5-diphenyltetratetrazolium (MTT) colorimetric assay. 1·10⁴ MCF-7 cells were seeded in a 96-well plate. To 100 μl of cells suspension 20 μl of MTT solution (5 mg/ml PBS, Sigma) was added. After that cells were incubated with MTT during 4 h in standard conditions. Mitochondrial dehydrogenases of viable cells cleave the tetrazolium ring, yielding purple MTT formazan crystals which are insoluble in aqueous solutions. The resulting purple solution is spectrophotometrically measured. An increase in cell number results in an increase in the amount of MTT formazan and an increase in absorbance. Then samples were centrifuged under 1500 g during 5 min and supernatant was extracted. In all wells were added 10 μl DMSO (Sigma) for MTT crystals dilution and 20 μl of 25 mM glycine. Optical absorption was detected on multi-well spectroscopy reader Multiscan (Labsystem, Finland) (OP540 nm).

**Statistical analysis.**

In common case of the biochemical studies a random variable is obviously normal or close to normal distribution, but a set of samples has a small amount that is not sufficiently representative. As with the basic parameters can not be estimated reliably, there is a need for statistical evaluation of selective parameters. Section of mathematical statistics, which is dedicated to the processing of small samples (2≤n<20), is known as micro-statistics. Underlying micro-statistic estimates of normally distributed random variables is the student's distribution for small quantities. Pearson correlation coefficient was used to determine the relationship between exposure and response to experimental biological systems and deficiencies of covariance. Coefficient was calculated for the cell viability in culture and the molecular mass of polyethylene glycol by the formula [19]:

\[
 r_{XY} = \frac{\text{cov}(X,Y)}{\sigma_X \sigma_Y} = \frac{\sum(X - \bar{X})(Y - \bar{Y})}{\sqrt{\sum(X - \bar{X})^2 \sum(Y - \bar{Y})^2}}.
\]

\[
 \bar{X} = \frac{1}{n} \sum_{i=1}^{n} X_i, \quad \bar{Y} = \frac{1}{n} \sum_{i=1}^{n} Y_i \quad \text{average of the samples}
\]

The correlation coefficient ranges from minus one to plus one. Thus for independent parameters it is 0, and for closely related approaches to the module of one. For statistical analysis processing of results and evaluation carried out by student’s test. Reported p values was *p≤0.05 or **p≤0.01.

**Table 2 – Cryopreservation solutions studied on the cell viability stage (end concentration)**

<table>
<thead>
<tr>
<th>Name</th>
<th>FBS, %</th>
<th>Saccharose, M</th>
<th>PEG(MM), %</th>
<th>DMSO, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>VS2.1</td>
<td>10</td>
<td>–</td>
<td>–</td>
<td>10</td>
</tr>
<tr>
<td>VS2.2</td>
<td>20</td>
<td>–</td>
<td>–</td>
<td>10</td>
</tr>
<tr>
<td>VS2.3</td>
<td>10</td>
<td>0.2</td>
<td>–</td>
<td>5</td>
</tr>
<tr>
<td>VS2.4</td>
<td>10</td>
<td>–</td>
<td>(1500) 10</td>
<td>5</td>
</tr>
<tr>
<td>VS2.5</td>
<td>10</td>
<td>–</td>
<td>(4000) 10</td>
<td>5</td>
</tr>
<tr>
<td>VS2.6</td>
<td>10</td>
<td>0.2</td>
<td>(1500) 10</td>
<td>5</td>
</tr>
<tr>
<td>VS2.7</td>
<td>10</td>
<td>0.2</td>
<td>(4000) 10</td>
<td>5</td>
</tr>
<tr>
<td>VS2.8</td>
<td>10</td>
<td>0.2</td>
<td>(15 000) 10</td>
<td>5</td>
</tr>
<tr>
<td>VS2.9</td>
<td>10</td>
<td>0.2</td>
<td>(20 000) 10</td>
<td>5</td>
</tr>
<tr>
<td>VS2.10</td>
<td>10</td>
<td>0.2</td>
<td>(40 000) 10</td>
<td>5</td>
</tr>
</tbody>
</table>
RESULTS AND DISCUSSION.

Cooling rate dependencies.

Cooling rate was determined by the area where the temperature graph was similar to linear: approximately between -10 °C and -190 °C.

As shown in Table 3, cooling rate of pure 0.2 M glucose solution (VS 1.1, Fig. 1 f) was bigger than pure 0.2 M saccharose solution (VS 1.2, Fig. 1 g) and other pure and mixed studied solutions (VS 1.3-1.10, Fig. 1 h-o). Obviously, reducing the concentration of CPA leaded to decrease of cooling rate. 5 % solutions of glycerol (VS 1.3) and DMSO (VS 1.5) demonstrated similar results, but 2.5 % glycerol showed better result, than the 2.5 % DMSO (VS 1.6). In mixed solutions samples with 5 % glycerol (VS 1.7, 1.10) were indicated better results, than for samples with 2.5 % glycerol and 2.5 % DMSO (VS 1.8, 1.9). Also, the mixed solutions with glucose (VS 1.9, 1.10) showed slightly better results, than the samples with saccharose (VS 1.7, 1.8), but the difference was negligible (Fig. 1).

No correlation between cooling rate and such parameters, as molar mass, molar concentration and concentration of polar groups, was observed at this stage of investigation. But correlation between molecular mass and cooling rate in the line of PEG was exist (Table 4). For the determination of the mechanism of dependency between cooling rate and composition of complex cryoprotectant solutions further researches are required.

Table 3 – Cooling speed in cryopreservation solutions

<table>
<thead>
<tr>
<th>Name</th>
<th>VS 1.1</th>
<th>VS 1.2</th>
<th>VS 1.3</th>
<th>VS 1.4</th>
<th>VS 1.5</th>
<th>VS 1.6</th>
<th>VS 1.7</th>
<th>VS 1.8</th>
<th>VS 1.9</th>
<th>VS 1.10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cooling rate, °C/min</td>
<td>322.0</td>
<td>261.3</td>
<td>279.6</td>
<td>274.3</td>
<td>281.3</td>
<td>192.0</td>
<td>276.7</td>
<td>235.3</td>
<td>257.0</td>
<td>279.6</td>
</tr>
</tbody>
</table>

Table 4 – Cooling speed in cryopreservation solutions with PEG

<table>
<thead>
<tr>
<th>Thermo couple in: MM, Da</th>
<th>Cooling rate, °C/min</th>
<th>From -4 °C to -40 °C</th>
<th>From -40 °C to -200 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>LN –</td>
<td></td>
<td>14 118</td>
<td>19 500</td>
</tr>
<tr>
<td>VS 2.6 – 1500</td>
<td></td>
<td>140.1</td>
<td>278.8</td>
</tr>
<tr>
<td>VS 2.7 – 4000</td>
<td></td>
<td>145.3</td>
<td>263.5</td>
</tr>
<tr>
<td>VS 2.8 – 15 000</td>
<td></td>
<td>150.2</td>
<td>258.1</td>
</tr>
<tr>
<td>VS 2.9 – 20 000</td>
<td></td>
<td>177.6</td>
<td>255.6</td>
</tr>
<tr>
<td>VS 2.10 – 40 000</td>
<td></td>
<td>180.3</td>
<td>131.4</td>
</tr>
<tr>
<td>Pearson' correlation</td>
<td></td>
<td>0.89</td>
<td>-0.92</td>
</tr>
</tbody>
</table>

Cell viability after cycle of freezing/thawing.

The survival percentage of the human somatic cells was determined by MTT test after 3 days of cultivation. In Table 5 and Fig. 2 was demonstrated how the composition of the vitrification medium affects on cell viability after freezing/thawing cycle.

According to obtained data the smallest cell survival after a cycle of freezing/thawing obviously observed in 10–20 % serum and 10 % DMSO (VS2.1, VS 2.2, Table 2, Table 5, Fig. 2).

Adding 200 mM of Saccharose (VS 2.3) to medium significantly increased the percentage of alive cells. Since VS 2.3 was used in the laboratory as a carrier solution, effectiveness of another solutions was compared only with VS 2.3 (Fig. 2). After following measurements of cell survival rate it was observed that the presence of 10 % of low molecular weight PEG (1.5 kDa and 4 kDa) in vitrification solutions lead to substantially increase the percentage of alive cells in culture after freezing/thawing cycle. For PEG 1.5 kDa cell viability was 132 % without saccharose and 196 % with saccharose in comparison with control. For PEG 4 kDa without saccharose 107.9 % viability was not statistically significant, but with saccharose it was more that 200 % of cell viability. Further increase of PEG molecular weight (15–40 kDa) was ineffective to the preservation of cell mass (Table 5, Fig. 2).

Our data also confirmed the results of cell cultures staining on 7-th day of cultivation after freezing/thawing cycle in the presence of various concentrations of cryoprotectants (Fig. 3).

So, we can conclude that the presence of high molecular compounds in vitrification solutions leads to smooth the curve of temperature decrease and to slower temperature decrease, compared to the
Fig. 1. Cooling rate of CPA solutions: a – FBS+DMEM; b – Saccharose 0.2 M; c – PEG 15; d – PEG 20; e – PEG 40; f – VS1.1; g – VS1.2; h – VS1.3; i – VS1.4; j – VS1.5; k – VS1.6; l – VS1.7; m – VS1.8; n – VS1.9; o – VS1.10.
saccharose solution. It slows the freezing speed and probably reduced cell survival rate. At the same time low molecular PEG (1.5 kDa) was more effective cryoprotectant, than high molecular PEG and saccharose. Adding of low molecular PEG to the vitrification solution increased the cell viability in 1.3 times (1.96–2.12 times with saccharose). Simultaneously, the presence of saccharose provided more effectively freezing conditions, that can be explained by several mechanisms. Firstly, presence of saccharose leads to osmotic dehydration of cells and reduction of intracellular ice formation. Secondly, saccharose provides the energy source to the cells for quick recovery after thawing that leads to increase of living cells percentage. Presence of PEG 1.5 kDa, according to the literature data, leads to increase of osmotic concentration of all components of the intracellular environment, suspending of biochemical processes in the cell, chromatin condensation [20] and modifying of the

Fig. 2. The viability of the cell culture after freezing/thawing cycle in the presence of various concentrations of cryoprotectants; * – p≤0.05; ** – p≤0.01; † – control.

Fig. 3. Cell cultures on 7-th day of cultivation after freezing/thawing cycle: a – VS 2.1; b – VS 2.2; c – VS 2.3; d – VS 2.4; e – VS 2.6; f – VS 9.9.
structural properties of vitrification of human somatic cells were analyzed. The possibility of increasing cells survival by adding polyethylene glycol of different molecular weights was shown. It was found that the low molecular weight PEG (1.5 kDa) exhibited a sufficiently high cooling rate (-140 °C/min) in the temperature range from -4 °C to -40 °C, and a high cooling speed (-278 °C/min) in the range -40 °C to -200 °C. Simultaneously, PEG (1.5 kDa) allowed to achieve the highest cell survival up to 200 % in comparison with control samples. Thus, the physico-chemical and molecular biological properties of low molecular weight PEG suggests the possibility of its application in vitrification environment for effective freezing protocols.

REFERENCES

ОПТИМІЗАЦІЯ ПРОТОКОЛІВ КРИЮЗБЕРЕЖЕННЯ СОМАТИЧНИХ КЛІТИН ЛЮДINI ЗА ДОПОМОГОЮ ПОЛІЕТИЛЕНГЛІКОЛЮ

Резюме
Було проаналізовано використання поліетиленгліколю при оптимізації протоколів кріозбереження соматичних клітин людини. Низькомолекулярний поліетиленгліколь як добавка до вітрифікаційних розчинів провив велику швидкість охолодження і продемонстрував виживаність клітін у 200 % порівняно з контрольною групою. Це дозволяє рекомендувати застосування низькомолекулярного поліетиленгліколю для вітрифікаційних середовищ для ефективних протоколів вітрифікації клітин.

КЛЮЧОВІ СЛОВА: кріозбереження, вітрифікація, поліетиленгліколь, швидкість охолодження, виживаність клітин, протоколи заморожування.

ОПТИМІЗАЦІЯ ПРОТОКОЛОВ КРИОСОХРАНЕНИЯ ЧЕЛОВЕЧЕСКИХ СОМАТИЧЕСКИХ КЛЕТОК С ПОМОЩЬЮ ПОЛИЭТИЛЕНГЛИКОЛА

Резюме
Было проанализировано использование полиэтиленгликоля при оптимизации протоколов криосохранения человеческих соматических клеток. Низкомолекулярный полиэтиленгликоль как добавка к витрификационным растворам проявил высокую скорость охлаждения и продемонстрировал выживаемость клеток в 200 % по сравнению с контрольной группой. Это позволяет рекомендовать применение низкомолекулярного полиэтиленгликоля для витрификационных сред для эффективных протоколов витрификации клеток.

КЛЮЧЕВЫЕ СЛОВА: криосохранение, витрификация, полиэтиленгликоль, скорость охлаждения, выживаемость клеток, протоколы замораживания.

Received 21.07.16

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