The effects of poly L-lactic acid nanofiber scaffold on mouse spermatogonial stem cell culture

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Introduction: A 3D-nanofiber scaffold acts in a similar way to the extracellular matrix (ECM)/basement membrane that enhances the proliferation and self-renewal of stem cells. The goal of the present study was to investigate the effects of a poly L-lactic acid (PLLA) nanofiber scaffold on frozen-thawed neonate mouse spermatogonial stem cells (SSCs) and testis tissues.

Methods: The isolated spermatogonial cells were divided into six culture groups: (1) fresh spermatogonial cells, (2) fresh spermatogonial cells seeded onto PLLA, (3) frozen-thawed spermatogonial cells, (4) frozen-thawed spermatogonial cells seeded onto PLLA, (5) spermatogonial cells obtained from frozen-thawed testis tissue, and (6) spermatogonial cells obtained from frozen-thawed testis tissue seeded onto PLLA. Spermatogonial cells and testis fragments were cryopreserved and cultured for 3 weeks. Cluster assay was performed during the culture. The presence of spermatogonial cells in the culture was determined by a reverse transcriptase polymerase chain reaction for spermatogonial markers (Oct4, GFRα-1, PLZF, Mvh/VASA, Igfα, and Igfβ1), as well as the ultrastructural study of cell clusters and SSCs transplantation to a recipient azoospermic mouse. The significance of the data was analyzed using the repeated measures and analysis of variance.

Results: The findings indicated that the spermatogonial cells seeded on PLLA significantly increased in vitro spermatogonial cell cluster formations in comparison with the control groups (culture of SSCs not seeded on PLLA) (P<0.001). The viability rate for the frozen cells after thawing was 63.00% ± 3.56%. This number decreased significantly (40.00% ± 0.82%) in spermatogonial cells obtained from the frozen-thawed testis tissue. Both groups, however, showed in vitro cluster formation. Although the expression of spermatogonial markers was maintained after 3 weeks of culture, there was a significant downregulation for some spermatogonial genes in the experimental groups compared with those of the control groups. Furthermore, transplantation assay and transmission electron microscopy studies suggested the presence of SSCs among the cultured cells.

Conclusion: Although PLLA can increase the in vitro cluster formation of neonate fresh and frozen-thawed spermatogonial cells, it may also cause them to differentiate during cultivation. The study therefore has implications for SSCs proliferation and germ cell differentiation in vitro.

Keywords: PLLA nanofibers, tissue cryopreservation, testis

Spermatogonial stem cells (SSCs) have the potential to self-renew and generate differentiated germ cells that will eventually lead to sperm.1 These cells can therefore play an important role in treating infertility, especially when it comes to cancer survivors who have been affected by the long-term adverse effects of cancer treatments:
To store and preserve mature sperm prior to treatment is common practice for those other than prepubertal cancer patients. In such patients, however, the preservation of the male germ-line cells is a challenge. It should be noted here that testis tissue cryopreservation and then proliferation and autotransplantation of isolated SSCs can facilitate the medical application of these cells.

The microenvironment of SSCs in the basal compartment of the seminiferous epithelium is important for the maintenance and self-renewal of these cells because Sertoli cells provide the growth factors necessary for self-renewal in this microenvironment. Glial cell line-derived neurotrophic factor (GDNF) is the most crucial factor in the balance between self-renewal and differentiation in the SSC pool as well as the promotion of SSCs’ self-renewal. Without this factor, either spermatogonial aggregates do not develop or SSCs perish. The generation process of SSCs and spermatogonia and also the localization of undifferentiated spermatogonia along specific portions of the basement membrane are done via stimuli from the vascular network and interstitial cells, ie, the peritubular myoid cells and the Leydig cells. On the other hand, the adhesion molecules of the basement membrane are anchored with SSCs.

Nanofiber matrices mimic the architecture and size scale of the natural extracellular matrix (ECM). This scaffold provides more three-dimensional (3D) topographical signals to seeded cells and results in a more physiologically relevant cellular phenotype compared with the two-dimensional substrates. It provides physical cues for cell orientation and spreading, and its pores make space for the remodeling of tissue structures.

Recently, many electrospun nanofibrillar surfaces have been used for cell culture. Poly L-lactic acid (PLLA) is one of the most promising biodegradable, biocompatible, and US Food and Drug Administration-approved polymers and can easily be electrospun to form a 3D non-woven network. To date, the proliferation of muscle-derived stem cells and the differentiation of hepatic cells from human mesenchymal stem cells have been established on PLLA. 3D soft agar culture system and electrospun polylamide nanofiber (Ultra-WebTM; Corning Life Sciences, Tewksbury, MA, USA) also have been reported to yield complete in vitro spermatogenesis of mouse testicular germ cells as well as short-term culture of spermatogonial stem-like cell colonies.

Culture and cryopreservation of SSCs are the most effective methods used for the long-term preservation of these cells. In addition, cryopreservation of cellular aggregates or tissues may allow for the isolation and culture of SSCs from frozen-thawed tissues in order to mimic conditions that will occur in oncology patients. To date, immature testicular pieces have been cryopreserved by slow-freezing and vitrification in both mice and humans and have been hetero-grafted beneath the tunica albuginea of a busulfan-treated recipient testis. These studies demonstrate that spermatogonia can survive in xenogeneic recipients after cryopreservation and therefore result in an offspring after sperm microinsemination. Our assumption here is that PLLA might provide an improved structural environment for the clonal expansion or differentiation of SSCs. While many efforts have been made to cryopreserve SSCs and testicular tissues in animals and humans, no report has yet been found with respect to the culture of SSCs obtained from frozen-thawed testis tissue on PLLA scaffolds.

For this reason, we have frozen and thawed immature testicular fragments following the removal of interstitial tissue with enzymatic digestion. We assumed that by combining the appropriate 3D scaffolds (provided by the biodegradable polymer scaffolds) with GDNF we could possibly create an appropriate environment for the proliferation of neonate mouse SSCs. The goal of the present study was to investigate the effects of a PLLA nanofiber scaffold on the frozen-thawed neonate mouse SSCs and testis tissue. We wanted to know whether in such a culture system, the SSCs would maintain the clonogenic and proliferation potential or instead differentiate.

**Materials and methods**

**Animals**

Eighty 3–6-day-old male mice from the National Medical Research Institute, initially from the original stocks of Razi Laboratory (Tehran, Iran), were used in the experiment. The animals were kept in plastic cages in a room maintained at a temperature range of 22°C–25°C, with a 12-hour light/dark cycle. The animals had free access to drinking water and standard laboratory pellets. The research was conducted in accordance with the National Research Council guidelines.

**Isolation and cultivation of spermatogonial cells**

Testes from the 3–6-day-old National Medical Research Institute mice were collected for the preparation of cell suspension following enzymatic digestions and purification steps. After decapsulation, the testes were minced and suspended in Dulbecco’s Modified Eagle Medium (DMEM; Life Technologies, Carlsbad, CA, USA) supplemented with 1.37 g/L NaHCO₃ (Sigma-Aldrich, St Louis, MO, USA) supplemented with 1.37 g/L NaHCO₃ (Sigma-Aldrich, St Louis, MO, USA).
USA), single-strength nonessential amino acids, penicillin (100 IU/mL), streptomycin (100 μg/m), and gentamycin (40 μg/mL) (all from Life Technologies). Testicular cells were separated by the method of van Pelt et al. with minor modifications. Briefly, minced testes pieces were suspended in DMEM containing 0.5 mg/mL collagenase/dispase, 0.5 mg/mL Trypsin, and 0.05 mg/mL DNAse, for 30 minutes (with shaking and a little pipetting) at 37°C. All enzymes were purchased from Sigma-Aldrich. For the next step, the interstitial cells were removed by washing in DMEM medium. A second digestion step was performed in DMEM media by adding a fresh enzyme solution into the seminiferous cord fragments as described above. After cell separation and filtration through 70-μm nylon filters, the collected cells were used for the culture cells. Immediately after cell isolation, the number of cells was determined using a hematocytometer. Viability was also assessed. Sertoli cells and myoid cells were also isolated through overnight differential plating in DMEM containing 10% fetal calf serum (FCS).

After the removal of the Sertoli and myoid cells, spermatogonia, which remained in suspension, were collected and cultured in DMEM containing 5% FCS and 10 ng/mL GDNF for 3 weeks. The cells were incubated at 32°C, 5% CO₂ in a humidified atmosphere, and the medium was refreshed three times per week. The diameters and the number of clusters were determined every 7 days during the culture for 3 weeks. The cells were subcultured during cell culture, and cluster assay was carried out on the 7th, 14th, and 21st days of the culture. Using an inverted microscope (Carl Zeiss; Oberkochen, Germany), the number of clusters and their diameters were measured by Image J software (version 1.240; National Institutes of Health, Bethesda, MD, USA). The identity of the cultured cells was confirmed by the expression of spermatogonial genes, ultrastructural study of cell clusters, and SSC transplantation to a recipient mouse.

**Cryopreservation and thawing procedure of SSCs and testis tissue**

The isolated cells and testis tissue fragmentations were cryopreserved using a procedure described by Izadyar et al. with some modification. Briefly, cell viability was assessed immediately after cell isolation. Cell suspensions in 0.5 mL aliquots (2 × 10⁶ cells per mL) were prepared. Then, an equal volume of 2× concentrated freezing medium was added dropwise to the Eppendorf vial containing the cell suspension, for a period of 10–15 minutes, and after gently mixing by inverting the vial, a sample was taken for viability assessment. The freezing media were based on DMEM supplemented with 10% (v/v) FCS, 1.4 M dimethyl sulfoxide (DMSO) and 0.07 M sucrose. Cryovials vials (1.8 mL; Nunc, Roskilde, Denmark) containing 1.0 mL of cell suspension in freezing medium were placed in −80°C for at least 1 day and then plunged into liquid nitrogen. The cells were thawed by swirling in a 38°C water bath for a period of 2 minutes. The contents of the vial was transferred to a tube and diluted slowly by adding two volumes, dropwise, of DMEM supplemented with 10% FCS. Then, the cells were pooled and centrifuged at 2000×g for 5 minutes, the supernatant was removed, and the pellet was resuspended in DMEM/FCS. A sample was taken for viability assessment, and the remainder of the cells was used for culture experiments.

For tissue cryopreservation, tubule fragmentations obtained from the first enzymatic digestion were transferred into a cryovial and cryopreservation solution was added in the same manner as the cell cryopreservation procedure.

**Fresh and cryopreserved spermatogonial cell culture on PLLA nanofibers**

A layer of PLLA nanofiber was used to provide an environment that resembled as closely as possible that of in vivo. PLLA nanofibers composed of PLLA and collagen fabricated by the electrospinning technique were purchased from Stem Cells Technology (Tehran, Iran). The PLLA nanofiber was used in a culture system with both cryopreserved and fresh SSCs. After placing the nanofibers on the dishes, fresh and frozen-thawed spermatogonial cells were seeded (5 × 10⁶ cells) on nanofiber and cultured in three groups: (1) fresh cells, (2) frozen-thawed cells, and (3) cells obtained from frozen-thawed testis tissue. In addition, fresh and frozen-thawed cells cultured on the plate without nanofibers were also considered as control groups. Cells were cultured for 3 weeks. The diameter and the number of colonies were determined every 7 days during the culture for 3 weeks. Cluster formation was assessed using the procedure described by Yeh et al.

**Identity confirmation of the spermatogonial cells**

**Ribonucleic acid (RNA) extraction and reverse transcription**

The presence of spermatogonial cells during the culture was determined by the expression of spermatogonial genes based upon previous animal studies. Total RNA from the 6-day-old testis tissue (positive control) and cultured testicular cells from the entire culture dish were extracted using a standard RNA extraction kit (Qiagen, Hilden, Germany) per the manufacturer’s instructions. The purity and integrity of the RNA was checked by a 260/280 nm ratio measurement.
In the reverse transcription reaction, 1 µg of total RNA was used with QuantiTect® Reverse Transcription Kit (Qiagen) per the manufacturer’s instructions.

Polymerase chain reaction (PCR)
The primers specific for promyelocytic leukemia zinc-finger (PLZF), octamer-binding transcription factor 4 (Oct4), GDNF family co-receptor α (GFRα-1), VASA homologue (Mvh), Itgal (β1-integrin), Itgα6 (α6-integrin), c-Kit, and GAPDH genes were designed using the previously described mouse sequences (Gene Bank) and Gene Runner software (version 3.02; Hastings Software Inc, New York, NY, USA) as shown in Table 1. GAPDH, a housekeeping gene, was included as an internal control to normalize the PCR reaction. Reverse-transcription PCR (RT-PCR) was performed using the prepared complementary deoxyribo-nucleic acid (cDNA), the primers, and with PCR Master Mix 2X kit (Fermentas, St. Leon-Rot, Germany) under the following conditions: 95°C for 3 minutes, followed by 35 cycles at 95°C for 30 seconds, under specific annealing temperature for each primer (PLZF, 55°C; Oct4, 60°C; GFRα-1.52°C; VASA, 62°C; Itgal, 52°C; Itgal, 55°C; c-Kit, 60°C; and GAPDH, 60°C) for 45 seconds, 72°C for 60 seconds, and a final extension of 72°C for 10 minutes. To separate PCR products, 1 µL of each sample was resolved on a 1.2% agarose gel, and electrophoresis was performed on a 1.2% agarose gel, and electrophoresis was performed using the Tris-Borate-EDTA (TBE) 1x loading buffer (Sigma-Aldrich), and the PCR products were analyzed on a 1.2% agarose gel, and electrophoresis was performed using the Tris-Borate-EDTA (TBE) 1x loading buffer (Sigma-Aldrich), and the PCR products were analyzed using UVItec software (version 12.6 for Windows; UVItec Ltd, Cambridge, UK). The ratios of the SSC-specific gene band intensities were compared with the corresponding GAPDH. All PCRs were independently replicated three times.

Quantifications of PCR products
Gene expression levels were examined by semi-quantitative RT-PCR. The PCR products were analyzed on a 1.2% agarose gel (Life Technologies BRL) and were visualized under ultraviolet transillumination after being stained with Gel Red. RT-PCR reaction was performed for SSCs and germ cell genes: PLZF, Oct4, GFRα-1, VASA, Itgal, Itgal, c-Kit, and GAPDH genes, and the intensity of each band was quantified using UVItec software (version 12.6 for Windows; UVItec Ltd, Cambridge, UK). The ratios of the SSC-specific gene band intensities were compared with the corresponding GAPDH. All PCRs were independently replicated three times.

Immunohistological staining
For immunohistological staining, the clusters were fixed with 4% paraformaldehyde (pH 7.4) for 20 minutes and were then twice washed with 0.1% Tween-20 in phosphate buffered saline (PBS) solution prior to blocking. The clusters were blocked with normal goat serum (Vector, Burlingame, CA, USA) in PBS for 15 minutes and were incubated overnight at 4°C with primary antibody solutions, including: rat polyclonal anti-α6-integrin (1:100; Sigma-Aldrich), rat polyclonal anti-β1-integrin (1:100; Sigma-Aldrich), rat polyclonal anti-Oct (1:100; Sigma-Aldrich), and mouse polyclonal anti-Thy-1 antibody (1:100, Santa Cruz Biotechnology, Santa Cruz, CA, USA). The next day, the clusters were washed with 0.1%

### Table 1 The sequence of the designed primers used for reverse transcriptase polymerase chain reaction

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sequence (5′–3′)</th>
<th>Annealing temperature (°C)</th>
<th>Size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mvh(VASA)</td>
<td>F.5′ GAT AAT CAT TTA GCA CAG CCT C 3′</td>
<td>59–60</td>
<td>149</td>
</tr>
<tr>
<td>Itgal</td>
<td>R.5′ GTC AAC AGA TGC AAA CAC AG 3′</td>
<td></td>
<td></td>
</tr>
<tr>
<td>c-Kit</td>
<td>R.5′ AAA CAC TAA TAG AGC CAG CA 3′</td>
<td>60</td>
<td>148</td>
</tr>
<tr>
<td>GFRα-1</td>
<td>F.5′ AAT TGT CTG CGT ATC TAC TGG 3′</td>
<td>60</td>
<td>130</td>
</tr>
<tr>
<td>Itgal</td>
<td>R.5′ ACA TCT GAT ATG AAG GGC AC 3′</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oct4</td>
<td>F.5′ GAC ATT ACT CAG ATC CAA CCA 3′</td>
<td>60</td>
<td>115</td>
</tr>
<tr>
<td>PLZF</td>
<td>R.5′ AGG TAG TAG AGA TCA ATA GGG AC T 3′</td>
<td></td>
<td></td>
</tr>
<tr>
<td>c-Kit/CD117</td>
<td>5′ CCCGTGGGGGTTGCAGCTAGAAA 3′</td>
<td>61</td>
<td>137</td>
</tr>
<tr>
<td>GAPDH</td>
<td>R.5′ CTGAAAGAGGCGGTTGCTAG 3′</td>
<td></td>
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</table>

Abbreviations: Mvh(VASA), mouse vasa-homologue; GFRα-1, GDNF family co-receptor α1; Itgal, integrin-α6; Itgal, integrin-β1; Oct4, octamer-binding transcription factor 4; PLZF, promyelocytic leukemia zinc finger; c-Kit, proto-oncogene c-Kit or tyrosine-protein kinase Kit; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; bp, base pairs.
Tween-20 in PBS (three times for 5 minutes) and incubated with respect to the secondary antibody: goat fluorescein isothiocyanate (FITC)-conjugated anti-mouse or anti-rat immunoglobulin G, (Sigma-Aldrich) diluted 1:200 for 45 minutes at room temperature. Next, the clusters were washed with 0.1% Tween-20 in PBS (three times for 5 minutes). Finally, all clusters were counterstained with 4,6-diamidino-2-phenylindole (DAPI) in PBS (Santa Cruz Biotechnology) diluted 1:2 for 10 seconds. Labeled cells were examined with a fluorescent microscope (BX51; Olympus), and images were acquired using an Olympus D70 camera. Negative controls were treated without the primary antibody.

**Ultrastructural study of cell clusters**

SSC clusters from groups were washed with PBS, pre-fixed in 2.5% glutaraldehyde in PBS (pH 7.4) for 2 hours, and post-fixed with 1% osmium tetroxide in the same buffer for another 2 hours. After dehydration in an ascending series of ethanol, specimens were placed in propylene oxide and embedded in Epon 812 (TAAB, Berkshire, UK). Semi-thin sections (0.5 mm) were stained with toluidine blue for light microscopy and ultrathin sections (60–80 nm) were contrasted with uranyl acetate and lead citrate before being examined by transmission electron microscopy (LEO 906; Carl Zeiss). The cluster cells were identified as primitive or differentiated spermatogonia based on cellular morphologies previously reported.46–48

**BrdU (5-bromo-2-deoxyuridine) cell labeling and transplantation into recipient mice**

Seventy-two hours before transplantation, BrdU was added to the medium in order to label and trace cells in recipient mice. This was done after the SSC clusters and the underlying somatic cells were trypsinized at the end of the 3 weeks.

Subsequently, the spermatogonial cells were transplanted into the seminiferous tubules of recipient mice via the rete testis that was treated with 35 mg/kg busulfan prior to transplantation. The treated recipient mice were devoid of endogenous spermatogenesis at the time of transplantation (6 weeks after treatment).49 Adult recipient mice were anesthetized with 10% ketamine and 2% xylazine (Alfasan, Woerden, The Netherlands). Approximately, 10^6 of the cultured cells in 10 µL DMEM were injected into the seminiferous tubules in one testis of each recipient mouse, while the other testis served as an internal control. Transplantation was performed by retrograde injection through the efferent ducts.50

**Statistical analysis**

The repeated measures and analysis of variance followed by Tukey post-hoc tests were used to evaluate differences between the experimental and control groups as well as between different time points in a group. Data are given as means ± standard deviation. The results were assumed significant when \( P \leq 0.05 \).

**Results**

**The effect of cryopreservation on the percentage of viable cells in experimental groups**

Cell viability was assessed after the isolation of testicular cells through enzymatic digestion. In this experiment, more than 89.25% ± 2.20% of the cells were determined viable in the fresh cell groups (control 1 and experimental 1). In the chemical toxicity test, however, adding freezing media did not seem to have a significant effect on viability, and more than 82.5% ± 4.2% remained viable. The viability rate of the frozen cells after thawing (control 2 and experimental 2) and spermatogonial cells obtained from frozen-thawed testis tissue (control 3 and experimental 3) were 63.00% ± 3.56% and 40.00% ± 0.82%, respectively. The viability rates decreased significantly in both groups when compared with the fresh cell groups (\( P \leq 0.001 \)).

**The effects of PLLA and cryopreservation on cluster formation of the SSCs in experimental groups**

The clusters, ie, 3D aggregations of germ cells on a feeder layer, appeared 2–3 days after the primary culture. These clusters were clumpy and had individually recognizable cells, and once enzymatically dispersed and re-plated, their SSC content could start new clusters during 3 weeks of culture (Figure 1A).

To determine the effects of PLLA on cluster formation of SSCs in vitro, experimental groups (culture of SSCs with seeding onto PLLA) were compared with their respective control groups (culture of SSCs without seeding on PLLA). Overall, the results indicated that during the 3-week cultivation, the number of clusters for all of the experimental groups significantly increased compared with their respective control group (\( P \leq 0.001 \)) (Figure 2A). Furthermore, a 3-week culture resulted in a significantly higher number of clusters compared with a 1- or 2-week cultivation (\( P < 0.05 \)).

To determine the effects of cryopreservation on the cluster formation of SSCs in vitro, a frozen-thawed cell group was compared with a fresh cell culture group. During the third week of culture, the number of clusters observed in the frozen-thawed cell group (control 2) as well as those in...
the cell group obtained from the frozen-thawed testis tissue (control 3) were significantly lower than that of the fresh cell group (control 1) (Figure 2A).

As shown in Figure 2B, differences in the diameter of clusters (µm) did not vary significantly from one week to another. Although the differences in the diameter of clusters (µm) in the fresh cell groups (both in the presence or absence of PLLA) were not significant, in the culture of only frozen-thawed cells onto PLLA (experimental 2), this diameter was significantly lower compared with the control group (P < 0.01) (Figure 2B).

Identity confirmation of the spermatogonial cells
RT-PCR and immunocytochemistry
To analyze the expression of specific spermatogonial and germ cell markers in the isolated testicular cells and cultured cells, RT-PCR was performed after 3 weeks of culture. As shown in Figure 1B, all samples expressed specific genes of spermatogonial cells (PLZF, Oct4, GFRα-1, VASA, Itgα6, and Itgβ1) and c-Kit as a differentiated germ cell gene (Figure 1B).

The presence of Itgα6 (Figure 3A–C), Itgβ1 (Figure 3G–I), Oct4 (Figure 3M–O) and thy-1 (Figure 3S–U) in cultured cells was confirmed by immunocytochemistry after 3 weeks of culture. Negative controls were treated without primary antibody (Figure 3D–F, J–L, P–R, and V–X).

Functional assay of the transplanted cluster cells
Germ cells were labeled with BrdU before transplantation. Immunofluorescence indicated that before transplantation, approximately 70% of cells had been labeled with BrdU (Figure 4A–C). After a month of cultivation, 10^5 cells from the third experimental group were injected into the seminiferous tubules through the rete testis of the recipient testes. This was done in order to confirm the presence of SSCs in clusters as well as to assess SSC colonization in the testes. One month after transplantation, those cells whose nuclei were positively stained with FITC for BrdU were considered the transplanted cells (Figure 4D–F). Transplantation studies revealed that compared with the non-transplant control group (Figure 4G–I), the transplanted cells localized as single cells only in the basal membrane of the seminiferous tubules of the recipient testes.

Ultrastructural characterization of cluster cells
The ultrastructural characteristics of spermatogonial cell clusters were examined via transmission electron microscopy. After 3 weeks of cultivation, the electron micrograph showed that cell clusters from the experimental culture groups (Figure 5A, D, and G) had the typical morphology of spermatogonial cells (Figure 5B, E, and H); however, cell differentiation was also observed among cell clusters obtained from the PLLA groups (Figure 5C, F, and I).
As shown in Figure 5B, E, and H, cell clusters had large spherical nuclei that contained one or two prominent nucleoli located either along the nuclear membrane or in the center of the nucleus. The shape of the cells was flat with either a long and regular or a round nucleus. In primitive spermatogonial cells, both the nucleus and cytoplasm are spherical. The spherical nucleus contains scattered flakes of heterochromatin. The nucleoli had a prominent reticulated nucleolome, and an irregular shape, and occupied an eccentric position in the nucleus. The cytoplasm was characterized by organelles, eg, mitochondria, rough endoplasmic reticulum, and vesicles that were mostly located in the perinuclear region. The mitochondria were clumped, interconnected by bars of cementing substance, and possessed parallel cristae. Spherical mitochondria were found single in relatively high numbers. The Golgi complex was poorly developed. In the differentiated spermatogonia, the nucleus was more heterochromatin; however, the shape of the heterochromatin was less dentate compared with the precursor cells.
Figure 4 Functional assay of spermatogonial stem cells into recipient mouse testis. (A) BrdU was added, and staining was examined in cultured spermatogonial cells before transplantation. (D) Transplanted cells on the base membrane of mouse seminiferous tubules 1 month after transplantation (arrowhead). These cells were traced in the recipient testes by BrdU staining. (G) The non-transplanted right testis was considered as the control group. (B, E and H) DAPI staining and (C, F and I) merge of FITC and DAPI.

Note: Scale bars = 200 µm.

Abbreviations: BrdU, 5-bromo-2-deoxyuridine; DAPI, 4,6-diamidino-2-phenylindole; FITC, fluorescein isothiocyanate.

Figure 5 Representative transmission electron micrographs from spermatogonial stem cell clusters (A–I). The electron micrograph showed cells from spermatogonial stem cell clusters in culture groups (A, D and G) on nanofiber scaffolds (Experimental 1–3) had morphology typical of spermatogonial cells. The nucleus shown contains a mottled appearance with dark speckles of heterochromatin (arrowhead; B, E and H). The cytoplasm was characterized by organelles, eg, mitochondria (M), rough endoplasmic reticulum (RER) and vesicles (Ve) that were mostly located in the perinuclear region. A portion of a small compact nucleolus (Nu) is visible and highly reticulated. In addition, the nucleus (N) of some cluster cells had marginal heterochromatin (Ht) and their morphology was similar to differentiated cells (C, F and I). The heterochromatin ratio increases in differentiated spermatogonial cells.

Note: Scale bar = 5 µm.
The effects of PLLA nanofiber scaffold on SSCs genes expression in culture

Gene expression of various genes was assessed and normalized based on GAPDH as a reference gene. Results showed lower gene expression of \( \text{Itg} \alpha 6, \text{Itg} \beta 1, \) and \( \text{Oct} 4 \) in the second experimental group compared with the second control group \((P < 0.05)\); however, no significant differences were observed in other gene expressions (PLZF, \( \text{GFR} \alpha 1, \), VASA, and c-Kit) among the groups (Figure 6). High expression of \( \text{Itg} \alpha 6 \) and \( \text{Itg} \beta 1 \) in the first experimental group was observed in the culture as opposed to the first control group \((P < 0.05)\). Also, gene expression of c-Kit in the fresh cell group significantly increased in comparison with the other genes.

Discussion

In this study, we were able to show that the culture of fresh or frozen-thawed SSCs as well as SSCs obtained from frozen-thawed testis tissue seeded on PLLA can increase the colony formation of SSCs in the culture system when compared with the control groups (without seeding the cells on PLLA). SSCs obtained from frozen-thawed testis tissue can also form many clusters after the freezing procedure in vitro.

In this study, we used 3–6-day-old mice testes, because in these animals, spermatogenesis begins immediately after birth.\(^{52}\) Only germ cells in the newborn mouse testis can be the gonocyte or spermatogonia, located in the center of the seminiferous tubule.\(^{53,54}\) By 6 days postpartum, these cells migrate to the basal membrane and produce undifferentiated type A spermatogonia, which begin to differentiate in a stepwise manner.\(^{55,56}\)

Since SSCs are rare in rodent testes (only 0.03% of all germ cells),\(^{57}\) and there are no clear surface specific markers for SSCs isolation, SSC culture and proliferation can provide new tools to investigate molecular mechanisms and signaling pathways that regulate SSC functions.\(^{19}\) In vitro culture may increase the rate of successful transplantation, even with a small number of dissected seminiferous tubules.\(^{56}\) One way to reach these goals may be to co-culture these cells within the presence of ECM components on 3D scaffolds. In this study, PLLA nanofiber scaffolds were used to mimic the structure of ECM. In recent years, PLLA has been widely explored as a biomaterial scaffold because of its impressive biocompatibility, biodegradability, minimal inflammatory reaction, and excellent mechanical properties.\(^{56}\) The electrospinning method has allowed the successful preparation

![Figure 6](https://www.dovepress.com/)

**Figure 6** Comparison between the relative gene expression of SSCs and germ cell genes in the experimental groups.

**Notes:** Values are mean ± standard deviation. The experiments were replicated at least three times. \(^*\)Significant difference versus culture of SSCs without seeding on PLLA in the same gene \((P < 0.05)\); \(^{**}\)significant difference versus all genes in the same group \((P < 0.05)\); \(^{**}\)significant difference versus fresh cells group in the same gene \((P < 0.05)\).

**Abbreviations:** Fresh, isolated testicular cells by two steps of enzymatic digestion before culture; Fresh+nano, fresh cells seeded on PLLA; Fz, frozen-thawed cells; Fz+nano, frozen-thawed cells seeded on PLLA; Fz+nano, frozen-thawed cells obtained from testis tissue; Fz+nano, frozen-thawed cells obtained from testis tissue seeded on PLLA groups; PLLA, poly-L-lactic acid; SSC, spermatogonial stem cell; Itg\( \alpha 6, \) integrin-\( \alpha 6 \), Itg\( \beta 1, \) integrin-\( \beta 1 \); VASA, mouse vasa-homologue; GFR\( \alpha 1, \) GDNF family co-receptor \( \alpha 1 \); Oct4, octamer-binding transcription factor 4; PLZF, promyelocytic leukemia zinc finger; c-Kit, proto-oncogene c-Kit or tyrosine-protein kinase Kit; GAPDH, glyceraldehyde-3-phosphate dehydrogenase.
of PLLA scaffolds with various topographical structures. The morphology and architecture of the electrospun structure is similar to that of the natural ECM. Indeed, randomly oriented PLLA nanofibers can enhance cellular infiltration within the in vivo scaffold.

To date, one established clinical option to preserve fertility is cryopreservation of sperm. There is no clinical technique currently available to preserve fertility in the prepubertal male who is likely to be sterilized by chemotherapy or radiotherapy, because he has no mature sperm. To overcome this problem, testicular tissues could be harvested before treatment and cryopreserved in these patients. Immature germ cells could somehow be matured either by autotransplantation, in vitro maturation, or xenografting.

In this study, the cryopreservation method of Izadyar et al was used to isolate neonate testicular cells as well as fragments of seminiferous tubules. Goossens et al cryopreserved mouse tissue pieces in ethylene glycol and DMSO as cryoprotectant while their protocols had already been used for freezing testicular stem cell suspensions by Frederickx et al. They obtained the best morphology of the basal compartment when the cryoprotective medium contained DMSO. Honaramooz et al investigated the effects of cooling or cryopreservation with DMSO on the testis fragments of pigs before grafting and observed complete spermatogenesis. Although some studies cryopreserved tissue pieces and successfully carried out grafting, they did not show isolated SSCs in order to increase cell number.

The survival rate of the frozen-thawed testicular cells (control 2) and the isolated cells after the freezing-thawing of the testicular tissue (control 3) were 63.00±3.56% and 40.00±0.82%, respectively, which decreased after cryopreservation in both groups. The SSCs that survived after cryopreservation were also able to form clusters in vitro. The survival rates of the frozen-thawed testicular cells obtained from our experiments were rather similar to the rates reported by Izadyar et al for prepubertal bovines (68%). Our study showed, however, that these rates decreased for those cells isolated after the freezing-thawing of the testicular tissue. Very likely, the decline may be the result of the diminished cell recovery following the freeze-thaw procedure. Previous studies have demonstrated that cryopreservation will preserve purified spermatogonial cells, because these cells are relatively resistant to freezing solutions compared with other spermatogenic cells. Other investigations have shown similar results for the cryopreservation of non-pure spermatogonia in other species including rodents and domestic animals as well as humans.

Since germ cell clusters have a distinct 3D structure, it may be possible to quantitatively analyze SSCs in vitro by counting clusters. In other words, by simply counting the number of colonies, the number of functional SSCs can be determined. In this study, we showed that the PLLA nanofiber significantly increased the number of clusters during the 3 weeks of cultivation. Additionally, the number of clusters in all groups significantly increased after the 3-week culture compared with a 1- or 2-week culture. Our finding is in line with Shakeri et al, who observed an increase of spermatogonial stem-like cell colonies during a short-term culture.

We also examined the effects of a PLLA nanofiber scaffold in combination with GDNF on the SSCs. Although several reports have described culturing various stem cells within the 3D scaffolds, the result of such cultures has most commonly been stem cell differentiation. Nur et al cultured embryonic stem cells on a 3D nanofibrous surface and observed proliferation with self-renewal. Because FGF-2 can promote the proliferation of embryonic stem cells, they also used FGF-2 with the nanofiber to proliferate these cells. Recent studies have shown that some soluble growth factors, especially GDNF, can have a long-term positive effect on SSC maintenance and may also stimulate SSC division in animals. We conclude that the presence of GDNF in combination with a nanofiber scaffold and the presence of somatic cells in the culture likely creates a testis-like microenvironment in which proliferation is promoted.

During cell culture, the number of clusters for both the frozen-thawed cell group and the cell group obtained from the frozen-thawed testis tissue were significantly lower compared with the fresh cell group. However, SSCs were probably more resistant to the freezing solution and the freezing-thawing procedure compared with Sertoli cells, which could not survive and died earlier in the procedure. Previous studies have demonstrated that somatic cells are able to differentiate or support SSCs in both mice and human cultures. Because Sertoli cells can secrete various growth factors or cytokines, they can therefore create a microenvironment that will promote the maintenance, survival, and proliferation of spermatogonia. We presume that the decline of the somatic cells may have reduced contact between Sertoli cells and SSCs in vitro and therefore decreased the chance of survival.
In spite of the decrease in cluster formation, SSCs may still remain alive in the culture without the ability to form clusters. This would be in accordance with our previous results that showed an increase in the gene expression of spermatogonial cells (Itgα6, Itgβ1, and Oct4) in frozen-thawed groups compared with the fresh cells. Previous studies have shown that B1-integrin is dominantly expressed in rodent SSCs and germ cells as a surface molecule. In this study, we conclude that spermatogonial cells are probably not as sensitive to the freezing–thawing procedure as are Sertoli cells, and so the number of these cells that can combine with different cell types that have no stem cell potential (Sertoli cells) are high in the culture system. In other words, the high expression of B1-integrin is possibly related to the presence of Sertoli cells in testsis.

In this study it was also demonstrated that gene expressions of spermatogonial cells decreased in cells cultured on PLLA. In contrast, the expression of the differentiation marker gene (c-Kit) increased in culture systems after the use of PLLA. Electron microscopy studies also confirmed the presence of differentiated cells in experimental groups. It is possible that the use of PLLA provides a structure for germ cells that allows for a disruption in the differentiation arrest of the spermatogonial cells in the culture system, and therefore cell differentiation is initiated in the experimental groups. Thus the expression of spermatogonial genes is decreased compared with the control groups.

Other studies have demonstrated that there are no specific biochemical or morphological markers for SSCs, but a combination of the expression of multiple markers can provide important information about spermatogonial cell types in rodents and other species. Therefore, in order to confirm the presence of spermatogonial cells during cultivation, an RT-PCR using spermatogonial and germ cells markers (Oct4, GFRα-1, PLZF, VASA, Itgα6, Itgβ1, and c-Kit) in all culture groups was performed. PLZF and GFRα-1 are markers for spermatogonial stem/progenitor cells and are well known spermatogonial-specific markers in many species and are considered to be markers of SSCs in rodents. Oct4 is a general marker for stem cells and is also expressed in mouse spermatogonial stem/progenitor cells. Itgα6 and Itgβ1 are cell surface receptors for spermatogonial stem/progenitor cells and mediate cell–cell and cell–ECM attachments and are expressed in germ cells of rodents. Mvh(VASA) as a marker of germ cell is expressed in all spermatogenic cells. The expression of c-Kit, the receptor for stem cell factor, is low or absent in A spermatogonia and enhanced in late A spermatogonia. In our study, the expression of Oct4, GFRα-1, PLZF, VASA, Itgα6 and Itgβ1 markers of spermatogonial and germ cell identification were observed in isolated testicular cells and all culture groups. Our findings are in line with results of previous research and support the aforementioned studies. Also c-Kit as a differentiated spermatogonia marker was observed in all culture groups, suggesting that the clusters contained differentiating germ cells. Kanatsu-Shinohara et al showed weak expressions for c-Kit in some colonies.

SSCs are the only cells capable of being recolonized in the seminiferous tubules of infertile animals. Therefore, cultured testicular cells were transplanted into a mouse busulfan azoospermic model through the efferent ducts, in order to examine the functionality of SSCs among the culture cells. A month later, colonization of spermatogonial cells in recipient mouse testes and SSCs was detected via BrdU staining. On the other hand, tubules with homing spermatogenesis were considered as colonized seminiferous tubules. In these tubules, spermatogonial cells were seen on the basal membrane of the cross-sectioned tubule of the recipient testes, which is in accordance with previous reports.

Moreover, an ultrastructure study of the cluster cells was performed. In rodents and humans alike, the relative amount of heterochromatin lining the nucleus is characteristic of specific types of spermatogonial cells. According to previous studies, the primitive spermatogonial group (A₁, A₆, and A₇) showed the large nucleus to cytoplasm ratio, intensive nucleolus, and low dense heterochromatin regions, whereas the heterochromatin face increased in differentiated spermatogonial group. Our findings are in line with previous studies.

Results of our study suggest that PLLA nanofiber scaffolds are useful in tissue engineering and can support the cell proliferation process. Frozen-thawed SSCs obtained from testis tissue can also form clusters after the freezing procedure in vitro. The self-renewal of SSCs in our cultural system enables this system to be utilized for the proliferation or differentiation of these cells in clinical applications, tissue engineering applications, cell replacement therapy, and tissue regeneration.

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Disclosure
The authors report no conflicts of interest in this work.

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