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Vulnerability of The Male Reproductive System to SARS-CoV-2 Invasion: Potential Role for The Endoplasmic Reticulum Chaperone Grp78/HSPA5/BiP

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Abstract

Severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) may adversely affect male reproductive tissues and male fertility. This concern is elicited by the higher susceptibility and mortality rate of men to the SARS-CoV-2 mediated coronavirus disease-19 (COVID-19), compared to the women. SARS-CoV-2 enters host cells after binding to a functional receptor named angiotensin-converting enzyme-2 (ACE2) and then replicates in the host cells and gets released into the plasma. SARS-CoVs use the endoplasmic reticulum (ER) as a site for viral protein synthesis and processing, as well as glucose-regulated protein 78 (Grp78) is a key ER chaperone involved in protein folding by preventing newly synthesized proteins from aggregation. Therefore, we analyzed Grp78 expression in various human organs, particularly male reproductive organs, using Broad Institute Cancer Cell Line Encyclopedia (CCLE), the Genotype-Tissue Expression (GTEx), and Human Protein Atlas online datasets. Grp78 is expressed in male reproductive tissues such as the testis, epididymis, prostate, and seminal vesicle. It can facilitate the coronavirus entry into the male reproductive tract, providing an opportunity for its replication. This link between the SARS-CoV-2 and the Grp78 protein could become a therapeutic target to mitigate its harmful effects on male fertility.

Keywords: COVID-19, Endoplasmic Reticulum, Grp78, Male Infertility, SARS-CoV-2

Introduction

The World Health Organization declared coronavirus disease 2019 (COVID-19), caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), as a global pandemic on March 11, 2020 (1). According to data from Johns Hopkins University’s Center for Systems Science and Engineering, there have been >400 million reported cases of COVID-19 worldwide, with >5 million deaths until July 2021 (https://coronavirus.jhu.edu/map. html) (2). Interestingly, clinical data emerging from patients demonstrate that male patients constitute 56-73% of the infected population (3). In addition, higher morbidity and mortality rates of SARS-CoV-2-infected males than age-matched females suggest sex-based differences in COVID-19 outcomes (4).

Severe acute respiratory syndrome coronavirus (SARS-CoV) and SARS-CoV-2 invade human cells through the angiotensin-converting enzyme-2 (ACE2) receptor and transmembrane serine protease 2 (TMPRSS2) (5, 6). As the 79% amino acid sequence identity of SARS-CoV-2 is similar to SARS-CoV (7), both viruses are thought to utilize the same receptor, ACE2, as a gate to mediate virus entry to target host cells (5, 8, 9). The higher affinity of SARS-COV-2 spike protein for binding to ACE2 (approximately 10 to 15 folds) compared to that of SARS-COV, which is one of the reasons for the more impactful pathogenicity of the latter (10). The subunit S1 of the S protein, which includes the receptor-binding domain, directly attaches to the peptidase domain of ACE2, while the membrane fusion is in control of the S2 subunit (11). In addition to ACE2, TMPRSS2, the cell-surface membrane protein, is required to integrate the virus with the cell membrane through cleaving SARS-COV spike proteins (10).

ACE2 is a metallopeptidase enzyme attached to the membranes of cells located in different
organs, particularly the lungs, heart, kidneys, and testes. Interestingly, ACE2 protein expression is evident only in specific tissues despite the demonstration of ACE2 mRNA in virtually all body organs (5). The physiological relevance of ACE2 in most tissues is yet unknown. However, ACE2 is regarded to be an important regulator of cardiac function and blood pressure control (12), probably through functioning as a natural counterpart to ACE1 (13). The testis is also one of the few organs with high levels of ACE2 expression (14, 15). ACE2 functions to convert Angiotensin II to Angiotensin 1-7 in Leydig cells and adjust testosterone production and consequently contribute to spermatogenesis modulation, which reveals their influence on male fertility (16). Overall, ACE2 mediates the activation of the Renin-Angiotensin-Aldosterone System (RAAS) (5) as a cell signaling system to regulate spermatogenesis (17).

Human spermatozoa include a number of RAAS family ligand enzymes and receptors, including the angiotensin receptor type 1 and 2 and the angiotensin mitochondrial assembly receptor (18). SARS-CoV-2 binding to this sperm surface signaling system may impair the functional capacity of the affected sperm. Specifically, the virus’s impact on spermatozoa ACE2 activity may enhance angiotensin II levels and increase reactive oxygen species (ROS) production resulting in oxidative stress (19). Excessive ROS production may cause sperm membrane and DNA damage and ultimately affect the sperm’s fertilizing potential (20).

Viruses such as human immunodeficiency virus (HIV), mumps, hepatitis B virus (HBV), influenza A virus subtype H1N1 (A/H1N1), and Ebola virus (21) had been implicated in the pathogenesis of orchitis, infertility, and testicular tumors (22, 23). Orchitis was also reported as a complication of SARS, an outbreak in 2003 caused by another member of the coronavirus family known as SARS-CoV-1. In another study, histopathological evaluation of testicular biopsy specimens from six men infected with SARS-CoV-1 indicated the findings of increased thickness of the basement membrane, destruction of germ cells, and low density of sperm in the seminiferous tubules (24). In this regard, a systematic review on the presence of SARS-CoV-2 in semen, including fourteen studies, suggested that the virus is rarely found in the semen of infected men and probably COVID-19 affects male fertility by making a deleterious effect on testicular structure (25). In line with this study, Khallili et al. (26) explained that despite the limited data on the detection of SARS-CoV-2 in the semen of infected patients, there is some evidence that the virus may play a role in testicular damage, abnormal sex hormone secretion, and infertility, which could be due to direct viral invasion through receptors or secondary immunological and inflammatory effects (26, 27). Hence, more studies are needed to evaluate all the possibilities.

The endoplasmic reticulum (ER) is the intracellular site where almost one-third of protein synthesis and protein folding occurs. Increased protein synthesis and excessive accumulation of unfolded/misfolded proteins in the ER lumen activate the unfolded protein response (UPR) and consequently drive the cell into ER stress (28). Under these circumstances, glucose-regulated protein 78 (Grp78), an ER chaperon protein, cooperates with three types of ER stress sensors such as Activating Transcription Factor 6 (ATF6), Protein Kinase R-like ER Kinase (PERK), and Inositol-requiring enzyme 1 (IRE1) to decrease unfolded/misfolded protein levels and avoid unfolded protein accumulation, thereby promoting cell survival. However, UPR may also activate the apoptotic response if ER homeostasis could not be regained (29). Therefore, Grp78 is more likely found within the ER lumen (30).

Under some circumstances, Grp78 can be moved towards the cell surface and act as a receptor to adjust different pathways (31). Grp78, also known as a binding immunoglobulin protein (BiP) or heat shock protein A5 (HSPA5) that is a member of the heat shock protein 70 (HSP70) family. This protein plays an essential role in resistance to apoptotic death in somatic cells and the response to chemical or physical cellular stress induced by cancer, malnutrition, and hypoxia (32, 33). Grp78 is highly expressed in human testicular tissue and mature spermatozoa, contributing to the physiology of spermatogenesis and fertilization (34, 35). Interestingly, previous results have confirmed that Grp78 protein contributes to the intrusion of different viruses like Ebola virus, dengue virus, influenza virus (36), Middle East respiratory syndrome coronavirus (MERS-CoV), Zika virus (37), and coronaviruses (38) into host cells. Moreover, recently the first experimental study proved that in addition to ACE2, the main receptor for virus entry, Grp78 as a host auxiliary factor for SAR-CoV-2 can simplify control virus entry (39).

Spike proteins of SARS-CoV-2 are one of the most important virulent factors of viruses to attach and penetrate host cells. For this purpose, host cell receptors like ACE2 and Grp78 are considered the target for viruses (5). Ibrahim et al. (40) have reported that the attachment of Grp78 Substrate Binding Domain β (SBD β) with the receptor-binding domain of the coronavirus spike protein is required to identify and help the virus to enter the host cells.

It has also been demonstrated that Grp78 protein expression increases in SARS-CoV infection, reflecting its role in virus entry into cells (41, 42). The present study aims to investigate the Grp78 expression in male reproductive organs using the findings of recent studies up to July 2021 and discuss the potential implications for SARS-CoV-2 invasion of the male reproductive tract.

In order to investigate the Grp78 expression in the male reproductive system, RNA and protein expression data of HSPA5/Grp78 in various human tissues and cancer, particularly male reproductive organs such as the testis and prostate were retrieved online using The Human Protein Atlas (http://www.proteinatlas.org/), Genotype-Tissue Expression (GTEx) (https://www.gtexportal.org), and
the Broad Institute Cancer Cell Line Encyclopedia (CCLE) (https://www.portals.broadinstitute.org/ccle) portals. All the data is available online. Protein expression scores are based on the best estimate of the “true” protein expression from a knowledge-based annotation. Immunohistochemistry (IHC) images of normal and cancer tissue of male reproductive organs such as testes, epididymis, and accessory sex glands from the tissue and pathology atlas on the Human Protein Atlas portal were used to evaluate the protein expression of HSPA5/Grp78 in the specific cells of these organs.

Data obtained from CCLE and the GTEx portal showed a high level of Grp78 mRNA expression in the male tissues such as the testis and prostate. The mRNA expression is also relatively high in the upper respiratory, digestive tracts, and lungs (Figs.1, 2). Data obtained from the Human Protein Atlas portal showed highly expressed in male tissues, including testis, epididymis, seminal vesicle, and prostate (Fig.3A, B). Moreover, IHC staining shows an increased level of Grp78 expression in cells of testis seminiferous tubules and the glandular cells of the epididymis, seminal vesicle, and prostate (Fig.3C). Data obtained from the Human Protein Atlas portal to assess Grp78 expression in various cancer organs, including the testis and prostate, showed high levels of this protein in these cancer organs (Fig.4).

Grp78 is overexpressed under pathological stresses like cancer, cellular malnutrition, hypoxia, and viral infections and is translocated from the ER to the plasma membrane (31, 32). This protein acts as a multifunctional receptor to interact with various proteins (29); therefore, it may be a gate for viruses to penetrate host cells (43, 44). In this regard, Grp78 has been introduced as a receptor to facilitate coronaviruses’ entrance into host cells in humans and bats (38). Thus, Grp78 seems to be an essential factor in helping virus protein folding, and its internalization to the host cells as well as protecting them from host immunity (Fig.5).

Recent studies proposed interaction between host cell Grp78 and the particular region of the COVID-19 spike model and considered this receptor a probable vaccination target (40, 45, 46). A series of recent studies have indicated that Pep42, a cyclic peptide, binds to the overexpressed Grp78 in the cancer cell membrane and by which enters the cancer cell (47, 48).
That is why Pep42 is considered a vehicle for tumor cell-specific chemotherapy (48). Ibrahim et al. (40) assessed the binding features of spike proteins of SARS-CoV-2 with Grp78. Interestingly, he also encountered 13 different cyclic regions in this spike model that were matched with the cyclic Pep42 structure and demonstrated the contribution of the spike protein model (regions III and IV) to binding with Grp78. Furthermore, a recent experimental study by Carlos et al. (39) revealed a new aspect of Grp78 role in the coronavirus infection. They presented some evidence that Grp78 functions not only as a cofactor to aid viral spike binding to ACE2, but also as a regulator of ACE2 protein expression, which highlight the great contribution of this protein to viral entry.

In the context of male reproduction, Grp78 expression has been confirmed by various studies in germ cells of humans and mice during spermatogenesis (49, 50). Investigation of Grp78 cellular localization in human testis revealed its expression in spermatocytes, round spermatids, and nick region of ejaculated spermatoozoa as well as principal cells of the epididymis (51). A similar observation was found using an animal model to measure the Grp78 gene expression in testicular tissue of two groups of 2-month and 4-month-old rats (52).

Based on the high level of Grp78 expression in male tissues, including spermatogenesis cells, epididymis cells, vesicle seminal, and prostate (Fig.3), we speculate that in addition to ACE2 and TMPRSS2, Grp78 can act as a receptor to intermediate coronavirus entrance to male reproductive cells. Leydig and Sertoli cells may be considered a target for SARS-CoV-2 due to the high expression of ACE-2 (8, 15). This may result in testicular destruction depending on the disease’s severity as immune and inflammatory responses (25). In this case, the virus attaches to the ACE2 receptor of the Sertoli cell and releases the viral RNA genome into the cytoplasm. The host ribosome translates the released RNA to produce fundamental viral protein and is finally inserted into the ER for processing (53).

On the other hand, several studies failed to demonstrate SARS-CoV-2 in the semen of COVID-19 patients, whether those patients were tested during an acute attack of the disease (54) or at different stages of recovery (55, 56). Pan et al. (55), and Stanley et al. (57), attributed the lack of SARS-Co-2 in the semen of COVID-19 patients to the fact that less than 1% of testis cells (spermatozoa, spermatogonia, and Leydig and Sertoli cells) express both ACE2 and TMPRSS2 receptors, which may reduce the virus’s ability to penetrate these cells. Accordingly, SARS-CoV-2 is unlikely to be sexually transmitted by men. However, it is difficult to rely on those observations to exclude the potential of sexual transmission of SARS-CoV-2 due to: i. Small sample of COVID-19 patients included in those studies and the heterogeneity of inclusion criteria, ii. Lack of data regarding the viral load of COVID-19- infected patients, iii. Difficulties...
that surround the collection of semen samples from COVID-19 patients during acute attacks, and iv. lack of a standard protocol of RT-PCR technique that is used for detection of seminal SARS-CoV-2.

The genome of SARS-CoV, like other coronaviruses, replicates in the host cell cytoplasm and is highly dependent on ER function for the preparation of proteins. This induces ER stress owing to the accumulation of unfolded yet synthesized SARS-CoV proteins in the ER lumen (28). Under these circumstances, UPR is activated by multiple cell-signaling pathways to maintain cellular homeostasis. However, if this condition continues and damages ER function severely, the UPR triggers cellular apoptosis (30). Interestingly, viruses apply various strategies to regulate UPR for ER preservation. In the case of SARS-CoV, UPR modulation is accomplished by the PERK pathway and eIF2α phosphorylation. This leads to the transcriptional activation of Grp78 as the intraluminal ER chaperones increase the processing and folding of expressed SARS-CoV proteins through viral replication and protect cells from apoptosis, at least in the early stage of infection (Fig.5) (41, 58).

Moreover, upon ER stress activation, IRE1α as an ER transmembrane sensor, starts generating X box-binding protein 1 (XBP1), acting as a transcriptional activator of genes involved in UPR to maintain ER and cellular function (30). It has been reported that SARS-CoV has been shown to cause a slight increase in XBP1 expression, which is likely important for increased virus protein folding and avoiding the harmful effects of ER stress-induced apoptosis (41). Therefore, SARS-CoV by selective modulation of the ER stress pathways provides the time and opportunity for viral replication before the infected host cell is sensitized to apoptosis.

It is notable that, in vitro treatment of lung epithelial cells with a humanized monoclonal antibody (hMAb159) with high affinity and specificity against GRP78, caused decreased cell surface GRP78 and cell surface ACE2 expression, as well as viral entry and SARS-CoV-2 infection. This finding showed that targeting host chaperones such as GRP78, which are necessary for viral entry and even production, might provide novel techniques for repressing SARS-CoV-2 and perhaps future coronavirus strains (39).

The connection between reproductive tissue cancers and increased Grp78 expression as a central part of UPR has been explored in prior studies. The results indicate a role of Grp78 in protein folding to guarantee survival, proliferation, and invasion of various cancer cells (59, 60) like the endometrial, gastric, renal cell, pancreatic, and prostate cancer (29, 61). These observations agree with our data extracted from the Human Protein Atlas portal (Fig.4). This may explain the relation of blocking Grp78 in different types of cancer cells and their apoptosis (62). Also, recently, some literature has proved the association between up-regulation and relocation of Grp78 to the tumor cell surface and some features like aggressive and invasive growth patterns of these cells (63, 64). This feature of cancer cells has turned the Grp78 at the cell surface into a useful prognostic marker and a target for cancer therapy (31).

A recent study by Liang et al. (65) concluded that patients with cancer have a higher risk of getting infected with SARS-CoV-2 compared to individuals without cancer. Although cancer patients are at a higher risk of getting infected as a result of immunocompromised states (66), the observation that Grp78 is overexpressed and translocated to the cancer cell membrane (67) may explain why tumor cells are more likely vulnerable to virus entrance and thereby virus propagation in cancer patients. Similarly, in diabetic and obese individuals, Grp78 is translocated to the cell membrane due to cellular glucose starvation (68), thus explaining why those individuals are at higher risk and severely affected (69). Based on these reports, increased Grp78 at the cell membrane may increase the severity of viral infection in specific tissue and individuals with higher Grp78 in their serum. Furthermore, it has been demonstrated that dysregulation of male hormonal sex can result from acute SARS-CoV-2 infection (70). Since Grp78 can play a prominent role in the steroidogenesis regulation of various reproductive mammalian cells (60). It seems that the hormonal defect is dependent on Grp78 function in testicular tissues infected with SARS-CoV-2.

**Conclusion**

In this study, we reviewed the present studies regarding the possible role of Grp78/HSPA5/BiP to facilitate the virus entry into host cells. However, it remains unclear how and when SARS-CoV-2 can impair male fertility potential. Given the global importance of fertility, all aspects of the impact of SARS-CoV-2 infection on the male reproductive system should be further assessed. A better understanding of the infection routes and target cells of the male reproductive system is critical for predicting some effective methods to treat or avoid likely consequences of infection in the male reproductive system.

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**Authors’ Contributions**

N.S.; Main search, collection, and study of published papers, writing, and approval of the manuscript. M.T., A.Sh., P.S., K.L., R.S., A.A., M.H.N.-E.; Reviewed the
manuscript, provided comments and suggestions, and finally approved the manuscript.

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Introduction

Successful fertility in mammals is dependent on various biological processes, including oocyte maturity, ovulation, embryo formation, and implantation (1). In majority of multicellular organisms, germ cells are the origin of new organism, which transfer the genetic and epigenetic information to the next generation. Furthermore, these cells are the main source of totipotency to create a new organism (2). Two important phases of gamete development are i. Primordial germ cell (PGC) formation during early embryogenesis and active migration to gonadal ridge and ii. Receiving distinct environmental signals for controlled cell meiosis division, in oogenesis and spermatogenesis processes (3). Considering the unique capabilities of PGCs in the production of gametes, these cells would be precious resources in reproductive biology and regenerative medicine. Thus, laboratory production of primordial germ cell-like cells (PGCLCs) has been a growing trend for years.

Stem cells can build specialized cells of human body, and exert self-renewal and differentiative capacity (4). In general, stem cells are categorized as “embryonic” and “adult”. Pluripotent embryonic stem cells (ESCs) can generate all cell lineages of human tissues through differentiation into more specialized multipotent stem cells with ectodermal, mesodermal, or endodermal origin. ESCs were first isolated from mouse embryo in 1981 (5). The pluripotent characteristics of ESCs have given treatment hope in patients suffering from different diseases including infertility (6). However, the use of ESCs has been associated with tumorigenesis and ethical concerns, and researchers have tried to replace these cells with adult stem cells (7, 8).

Adult stem cells, also known as “somatic” stem cells, are found both in developing and adult tissues (9). Mesenchymal stem cells (MSCs) are multipotent stromal cells which are highly found in adipose tissue, umbilical cord blood and bone marrow; however, they can also be identified in other tissues and regions requiring wound healing (10). MSCs can be differentiated into different cell types with mesodermal and non-mesodermal origin such as adipocytes, osteoblasts, and chondrocytes (11). The easy harvesting of some MSCs types, including umbilical or adipose tissue derived MSCs, immunosuppressive properties, fewer ethical concerns, as well as simple and cost-effective culture and differentiation methods have made them a suitable alternative for ESCs and induced pluripotent stem cells (iPSCs) (12).

Remarkably, PGCLCs can be differentiated not only from ESCs, but also reprogrammed pluripotent stem cells (PSCs) generated either through somatic cell nuclear transfer or induced pluripotency (13). Differentiation induction in transdifferentiated cells is a newer strategy for in vitro production of PGCLCs. iPSCs are obtained by reprogramming mature somatic cellsthat have been used successfully to produce PGCLCs (14).
transdifferentiation process, somatic stem cells are reprogrammed into cells of other germ layers and tissues, which is characterized by losing germ layer properties. The process of transdifferentiation to PGCLCs has been successfully induced in several somatic stem cell types such as skin-derived stem cells (15). The application of reprogramming and transdifferentiation prospectively circumvents the strict ethical limitations associated with obtaining PSCs from human embryos. In addition, genetically modified gametes can be obtained eventually using gene editing. However, somatic cells have rather different pattern of mutations (16) and epigenetic status (17) than that of germ cells. These differences can only be identified and characterized by strict monitoring at the genetic level, which have not yet been well developed.

Induction of PSCs into PGCLCs has been examined in two-dimensional culture (18) and embryoid bodies (19). Despite inherent differences, in both mouse and human PSC types, bone morphogenetic proteins (BMPs) have been identified as essential inducers of PGC specification. Using cytokines such as retinoic acid, co-culture with somatic cells and conditioned media, successful experiments have been performed. Further, in other studies by manipulation of transcriptional regulators expression deleted in azoospermia-like (Dazl) family genes (20) or using small molecules such as kinase inhibitors, successful PGCLC meiosis induction have been achieved in human iPSCs (21).

Wnt signaling plays an important role in gastrulation process especially in mesoderm and endoderm differentiation. It has also been suggested that Wnt signaling inhibition stabilizes the undifferentiated state of PSCs. Moreover, during induction of human iPSCs in defined conditions through Wnt signaling, an initial differentiation stage to mesoderm-like cells has been identified (14). These cells express genes such as Eomesodermin promoting commitment to PGCLCs of iPSCs through Sox17 upregulation (22). Such a phenomenon has also been shown in PSCS (23). Subsequently, using BMP4, mesoderm-like cells were differentiated into PGCLC (24). In this process, B lymphocyte-induced maturation protein 1 (Blimp1) suppresses the "neuronal differentiation" program and its expression is as a key feature of the PGC specification (14). Shirzeily et al. (25) in their study also demonstrated the differentiation ability of mouse adipose tissue and bone marrow-derived MSCs into primordial germ cells by expressing Mvh, Dazl, Stra8, and Scp3 specific markers.

Based on the findings, it is hypothesized that the mesoderm-like cells might be efficient precursors to form germ cell line with fewer ethical considerations than ESCs and iPSCs (26).

This review aims to overview the utilized MSC types and differentiation protocols for in vitro germ cell production with a focus on human and certain other mammalian models. Initially, the background of PGC development in the mouse model will be reviewed. Then, the existing studies on the production of PGCLCs from MSCs will be discussed by cell origin. Finally, the relevant information on all-trans-retinoic acid (RA) and BMP4 participation, as common factors used in the PGC specification, will be outlined.

The search for published records was carried out in the PubMed, EMBASE (Elsevier) and Google Scholar in January and August 2021 without limiting the search by date of publication and geographical region. The search terms were "mesenchymal stem cell", "mesenchymal stromal cells", "Wharton Jelly cells", "mesenchymal progenitor cells", "germ cells", "oogonial stem cells", "germline stem cells", "primordial germ cells", and "primordial germ cell-like cells". In addition, the operators "AND" and "OR" were applied for "primordial germ cells" or "primordial germ cell-like cells" and the other terms.

Authors independently screened the records by reading the title and abstract. Only peer reviewed full-length records covering the mammalian MSCs differentiation into primordial germ cell-like cells were included by experimental models.

In initial search, we retrieved 159 potentially relevant records, from which 50 were duplicates. Of the papers screened by reading the title and abstract, 77 studies were included (Fig.1).

![Fig.1: Flow chart of the review records selection.](image)

**Overview of mice primordial germ cell generation**

Wnt3A from trophoblasts together with BMP4 and BMP8b from amniotic adjacent mesenchyme inducing PGC specification. PGCs are the first population of germ cells which are established during the development (Fig.2) and are the immediate precursors of both oocytes and spermatogonia. In mice, PGCs are initially identified as an approximately 40-cell cluster in the incipient allantois
MSCs Differentiation into PGCLCs

based on day 7.25 (E7.25) of embryonic life. These cells then migrate to the developing hindgut endoderm and mesentery at E7.75 and E9.5, and colonize the genital ridges at E10.5, respectively (2).

An important event during the proliferative phase of PGCs is epigenetic reprogramming (27), especially a genome-wide DNA demethylation (28). Blimp1 and Prdm14 promote PGC specification via repressing somatic genes. PGCs retain remarkable pluripotent capabilities as shown by the ability to generate teratomas and pluripotent cell lines. Despite partial differences, both mouse and human PGCs express a group of ‘naive’ and ‘general’ pluripotency factors. A transcription program involving the expression of the RNA-binding protein Dazl subsides PGCs pluripotential capabilities and prime them toward germline commitment (29).

In female XX embryos, PGCs will continue the proliferation until E13.5 to reach a 25000 cell population. Afterwards, they enter to the prophase I of meiotic division and will be arrested in diploten in the spermatogonial stage of meiosis prophase I. In contrast, XY PGCs enter the mitotic arrest upon entry into the genital ridges, and stay silent in the G0/G1 phase of the cell cycle for the remaining embryonic period. Around day 5 postpartum many of these cells resume proliferation, whereby some of them will be recruited as spermatogonial stem cells (30).

Strategies for germ cell differentiation from mesenchymal stem cells

As summarized in Table 1, several studies have been reported that MSCs originally show germ cells characteristics, and in the presence of certain chemicals, they can also be differentiated into PGCLCs, potentially applicable as a therapeutic approach for infertility (31-33).

<p>| Table 1: Studies using mesenchymal stem cells to produce germ cell-like cells |
|-------------------------------|-----------------|------------------------------|-----------------|-----------------|-----------------|</p>
<table>
<thead>
<tr>
<th>Cell source</th>
<th>Cytokines</th>
<th>Additional strategy</th>
<th>Differentiation time (day)</th>
<th>Final cell type</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>BM-MSC</td>
<td>RA (µM)</td>
<td>BMP4 (ng/ml)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rat</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>21</td>
<td>PGCLC (11)</td>
</tr>
<tr>
<td>Mouse</td>
<td>-</td>
<td>20</td>
<td>-</td>
<td>4</td>
<td>PGCLC (32)</td>
</tr>
<tr>
<td>Mouse</td>
<td>-</td>
<td>1-25</td>
<td>-</td>
<td>4</td>
<td>PGCLC (33)</td>
</tr>
<tr>
<td>Mouse</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>2</td>
<td>PGCLC (25)</td>
</tr>
<tr>
<td>Ram</td>
<td>-</td>
<td>100</td>
<td>TGF-β1 treatment</td>
<td>21</td>
<td>Male GCLC (34)</td>
</tr>
<tr>
<td>WJ-MSC</td>
<td>Human</td>
<td>Co-culture with placental cells</td>
<td>14</td>
<td>PGCLC</td>
<td>(35)</td>
</tr>
<tr>
<td>Human</td>
<td>1</td>
<td>10</td>
<td>-</td>
<td>21</td>
<td>PGCLC (36)</td>
</tr>
<tr>
<td>Human</td>
<td>10</td>
<td>-</td>
<td>-</td>
<td>14</td>
<td>PGCLC (37)</td>
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<tr>
<td>Human</td>
<td>-</td>
<td>12.5</td>
<td>Overexpressed CD61</td>
<td>7</td>
<td>Male GCLC (38)</td>
</tr>
<tr>
<td>Human</td>
<td>-</td>
<td>-</td>
<td>Follicular fluid, cumulus cells conditioned medium</td>
<td>21</td>
<td>PGCLC (39)</td>
</tr>
<tr>
<td>Human</td>
<td>2</td>
<td>-</td>
<td>-</td>
<td>14</td>
<td>PGCLC (40)</td>
</tr>
<tr>
<td>Ad-MSC</td>
<td>Mouse</td>
<td>Co-culture with Sertoli cells</td>
<td>2</td>
<td>PGCLC (25)</td>
<td></td>
</tr>
<tr>
<td>Human</td>
<td>10</td>
<td>-</td>
<td>-</td>
<td>21</td>
<td>PGCLC (41)</td>
</tr>
<tr>
<td>Human</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>7-21</td>
<td>Male GCLC (42)</td>
</tr>
<tr>
<td>Human</td>
<td>10</td>
<td>-</td>
<td>Co-culture with Sertoli cells</td>
<td>21</td>
<td>Male GCLC (43)</td>
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<tr>
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<td>-</td>
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<td>Overexpressed miR-106b</td>
<td>4</td>
<td>PGCLC (44)</td>
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<td>-</td>
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<td>2 and 10</td>
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<td>BMP15 treatment</td>
<td>21</td>
<td>Female GCLC</td>
<td>(46)</td>
</tr>
</tbody>
</table>

Bone marrow-derived mesenchymal stem cells

BMP4 is one of the most frequently used factors for stem cell differentiation (47). Using BMP4, several protocols have been established to differentiate MSCs, including PGCLCs from bone marrow-derived (BM)-MSCs. Leading studies have used feeder layer, respectively and fetal bovine serum to support cell proliferation. For example, the mouse myoblast cell line C2C12 feeder layer in the presence of an interleukin 6 class cytokine was used to support proliferation (32). The cells obtained from rat, mouse, and human have been examined and various BMP4 concentrations have been employed from 1 to 100 ng/ml. In one study, RA was used as the PGCLCs differentiation inducer (11).

In the study by Ghasemzadeh-Hasankolaei et al. (34), although transforming growth factor (TGF)-β1 was more effective than BMP4, BMP4 could significantly boost the male germ cell markers. In addition to inducing differentiation, BMP4 had a positive effect on cell proliferation and survival (33). In three studies, the induction period was shorter than 5 days (25, 31, 33), while, in two studies the induction period was 21 days (11, 34). In two studies, mouse vasa homologue (Mvh) marker was employed to determine the extent of differentiation into germ cells (32, 33). This marker is expressed in germ cells up to the post-meiotic stage in both males and females (48).

In only two studies, isolated MSCs were characterized before induction of differentiation (11, 33). For this purpose, CD90+, CD105+, CD34 and CD45- molecular pattern was used to isolate and detect the cells. After inducing differentiation, several germ cell markers including c-Kit, Dazl, Stella, and Fragilis (also known as interferon induced transmembrane protein 3) were assessed. Fragilis is highly expressed in mouse PGCs at E6.5-7.25 together with Stella (Fig.2). In two studies, serum concentrations of 20% (32) and 10% (11) were used to support cell growth and proliferation. In the study of Shirazi et al. (32), comparison of migrating cells with migrated cells revealed a pattern of differentiation markers similar to PGCLCs. In one study the differentiation capabilities of BM-MSCs and ADSCs were compared. While both cell populations had the potential to become PGCLCs, BM-MSCs indicating a greater potential (25).

Wharton’s jelly-derived mesenchymal stem cells

Since BM-MSCs isolation is an invasive and complicated process, the use of umbilical cord or Wharton’s jelly (WJ)-MSCs is considered more convenient. The isolation of these cells from umbilical cord or placenta of a newborn is non-complicated, and non-invasive with a lower risk of contamination and ethical concerns (35). In addition to multipotent properties of these cells, which are between adult cells and ESCs, they seem to have a high potential to be differentiated into germ cells (49).

The strategies used to differentiate PGCLCs from WJ-MSCs are more diverse than BM-MSCs. Co-culture and genetic manipulation methods have been employed as supporting or adjuvant factors with RA and BMP4 treatments.

Fig.2: Female mice primordial germ cell (PGC) generation. The extraembryonic endoderm (ExE) layer, which is in contact with epiblast, secretes BMP4. Signals generated by ExE and visceral endoderm (VE) play important roles in inducing differentiation to PGCs. B lymphocyte-induced maturation protein 1 (Blimp1) is the first expressed marker of PGCs precursor at embryonic day (E) 6.25. BMP4 and BMP8b secreted by ExE and BMP2 secreted by proximal VE induce differentiation to Blimp1+ PGC precursors in the posterior proximal epiblast at E6.25. Initial PGCs at E6.5 and E7 express the transcription factors Prdm14 and Stella, respectively. By the fate determination of the germ cells, they begin to proliferate and migrate to the base of the yolk sac and show strong alkaline phosphatase (AP) activity. Then, they start migrating from inner cell mass (ICM) to the genital ridge. These cells express the pluripotency gene markers (PM) Oct4, Nanog, and Sox2, which are important for PGC growth and germination. Deleted in azoospermia-like (Dazl), mouse vasa homologue (Mvh) and ATP-dependent RNA helicase Ddx4 expression lead to germ cell commitment and indicate formation of post-migrating gonocytes. PGCs express the signaling by retinoic acid 8 (Stra8) from E13.5 which initiates the transition between mitosis and meiosis in oogenesis (26).

Compared with RA induction, the placental co-culture method further increased the early germ cell markers, Oct4, and ATP-dependent RNA helicase Ddx4. However, no significant differences were observed for specific cell markers such as growth differentiation factor 9 (GDF9) and zona pellucida glycoprotein 3 (Zp-3). GDF9 is a growth factor from the TGF-β family. This factor is expressed in large amounts in eggs that play an important role in folliculogenesis plus ovulation (50). Zp-3 as a receptor mediates the initial binding of sperm to the egg (51).

Based on these results, it seems that in MSCs differentiation induction into PGCLCs, BMP4 is not a substitutive factor, but a complementary or amplifying factor of RA. In support of this hypothesis, initial treatment
of human WJ-MSC with BMP4 and RA outperformed BMP4 alone. In addition, co-culture with placental cells and RA also showed fewer germ cell-specific markers than BMP4 and RA combination (36).

Human follicular fluid and cumulus cells-conditioned medium could induce expression of oocyte specific genes and proteins (39). These factors also induced morphological changes matching-oocyte-like cell differentiation. The observed effect has been related to potent growth factors of cumulus cells secretome such as epidermal growth factor. The potential of cells derived from follicular fluid to differentiate into oocyte-like cells further demonstrate the possibility of developing germ cells from adult stem cells (46).

In optimizing the development of PGCLCs, genetic manipulation is a potential strategy. Through overexpression and suppression of selected genes, the differentiation pathway can be oriented in the relevant direction. This approach can also facilitate in vitro induction of differentiation into PGCLCs. For this purpose, the genes involved in embryonic development are prioritized. In a study, obtaining male PGCLCs, CD61 or Integrin beta-3 was overexpressed in human placental MSCs. This manipulation alone increased the PGCLC markers such as c-Kit, sex determining region Y-Box 2 (Sry), and SSEA1. Treatment of manipulated cells with BMP4 enhanced male-PGCLC differentiation, which was characterized by an increase in the signaling through retinoic acid 8 (Stra8) marker (38).

Adipose-derived stem cells

It has recently been shown that MSCs derived from adipose tissue can be differentiated into PGCLCs with BMP4 treatment or transfection by miR-106b (44). BMP4 has not been used in any of the other five studies on adipose-derived stem cells (ADSCs). Nevertheless, RA has been used alone or in combination with co-culture. Genetic manipulation of ADSCs has also been described as a successful means to induce PGC specification.

The Sertoli cell co-culture (43), testicular cell-conditioned medium (42), and testosterone (43) were used to induce male PGCLCs with RA. In addition to direct differentiation induction, treatment with RA and testosterone indirectly enhanced the differentiation of ADSCs into male PGCLC by promoting the viability and secretory activity of sertoli feeder cells. Similar to human WJ-MSCs (48), increased CD61 expression in Canine ADSCs alone significantly elevated PGC specification and stem cell markers compared to control cells (42). Examination of TGF-β signaling showed that CD61 expression significantly enhanced the level of Smad2 phosphorylation, without affecting the level of phosphorylated Smad2/3 and Smad3.

Retinoic acid and bone morphogenetic proteins pathways

All-trans-retinoic acid

Retinol is obtained as retinyl ester from plants β-carotene. RA is one of the main retinol metabolites with potent biological capabilities related to proliferation and differentiation (52). On the cell surface, RA is taken up by retinol binding protein encoded by steroidogenic acute regulatory protein (StAR) 6 (53). Lecithin:retinol acyl transferase is also required for retinol uptake and esterification (Fig.3A). Inside the cell, the transferase and dehydrogenase enzymes convert retinol to retinyl ester, retinaldehyde, and then RA. RA binds to cellular RA binding protein 2 that is transferred into the nucleus (54). In the presence of RA, the retinoid-X receptor (RXR)/RA receptor (RAR) heterodimer complex interacts with DNA and activates the transcription of RA “primary response” genes (55).

Transcription activation is one of the primary steps of RA-associated differentiation process occuring during several minutes to hours after RA addition to the culture media. A number of “immediate early” genes or “primary response” genes are the direct targets of RA (Fig.3B) (56).

RA-mediated gene expression regulation often involves polycomb group (PcG) proteins (Fig.3C). PcG proteins can form a complex of gene-silencing proteins, which play a central role in embryogenesis, patterning, and differentiation (57). Following Ra in addition to stem cell culture medium, a fast dissociation of PcG proteins occurred, leading to the induction of differentiation-related genes expression, including Stella, Fragilis, and Stra8 (58). Thus, retinoids provide an essential early signal to induce a certain cascade for totipotent and lineage restricted stem cell differentiation (55).

**Fig.3:** Retinoic acid signaling pathway in the regulation of germ-cell related gene expression. A. On the cell surface, retinol (Rol) by binding to retinol binding protein enters the target stem cell and is converted to all-trans retinoic acid (RA) following the lecithin:retinol acyl transferase (LRAT), retinol dehydrogenase 10 (RDH10) and aldehyde dehydrogenase 1α2 (ALDH1A2) reactions. Then, RA is transferred to the nucleus by binding to cellular retinoic acid binding protein 2 (CRABP2). Once in the nucleus, RA is initially bound to retinoic acid receptors (RARs) and the RA-RARs complex by binding to retinoid X receptors (RXRs) will interact with RA primary response genes (53). B. RA primary response genes have some enhancers known as RA response element (RARE) which RARs/RXRs complex binds to, leading to RNPoi II activation and increased PGCs-related genes, including Stella, Fragilis, and Stra8, expression (55). C. PcG proteins form a gene-silencing complex for gene expression regulation. RA causes the dissociation of these proteins and activates the differentiation-related genes expression (57, 59).
bone morphogenetic proteins

PGCLC production from MSCs has been studied in both human and mouse models. Studies on the mouse and rat models have predominantly applied BM-MSCs. Despite promising early results on the differentiation of mouse BM-MSCs into PGCLCs, no follow-up data has been published on human BM-MSCs. MSCs of WJ and adipose tissue origin used in the human model studies (Table 1). However, the current knowledge on the mechanism of MSCs differentiation into PGCLCs is not persuasive enough. As the isolation of MSCs in humans is far easier than the obtaining ESCs, establishing suitable conditions for them to differentiate into PGCLCs will progress more rapidly. The recent studies that have efficiently differentiated MSCs are good models for future mechanistic studies, though failure to control the key variables remains a major limitation. Figure 4 illustrates the BMPs signaling pathway in the regulation of germ-cell related gene expression.

In vitro differentiation of MSC using RA and BMP4 has provided a customizable approach for improving conditions (Fig.5).

The available methods in this area have a number of limitations. Defining standard functional assays in this area will enable us to improve the conditions for producing PGCLCs. Evidence suggests that the mouse PSC is not very similar to the human PSC in terms of its pluripotency nature, making the PGCLC properties obtained from these two species different. For example, in very similar differentiation induction protocols, human PGCLCs, unlike mice PGCLCs, are negative for Ddx4 and Dazl genes, analogous to early stage PGCs (14, 61). Refined germ cell-related signaling pathways will enhance preferential differentiation of somatic MSCs into PGCLCs.

Discovering gene expression signatures using RNA sequencing methods will allow determining the functional potential of PGCLCs without the need for further differentiation. In determining such characteristics, the intrinsic differences between cell types from different species must be taken into account. The difference in the initial pluripotency state is attributed to the limited response of rat ESCs compared to mice ESCs to PGCLCs differentiation stimuli. Indeed, induced expression of the PGC transcriptional repressor Blimp1 by genetic manipulation increased differentiation towards cells expressing PGCLC markers (62). Moreover, human PGCs, unlike mice PGCs, do not express the pluripotency factor Sox2, while requiring Sox17 for PGC specification (24). Thus, depending on the studied species, additional strategies may be needed to obtain more mature PGCLCs, such as co-culture with somatic gonadal cells.

A reasonable similarity between ESC-induced PGCLCs and in vivo PGCs has previously been reported (63). However, no study has yet compared the transcript of PGCLCs obtained from MSCs with in vivo PGCs or PGCLCs from ESCs, reprogrammed cells, or transdifferentiated cells. The use of a CRISPR/Cas9 screening platform to identify the factors required for PGC development in a new study (64) is a good example of a mechanistic study of the PGC specification.

The development of safe and efficient protocols in this field will provide useful resources for the PGCLCs in fertility science. Certainly, a fully defined condition cannot be provided with somatic cell co-culture strategy
or serum supplementation. Nevertheless, using co-culture techniques, signaling and environmental cues related to PGC specification can be investigated. With these techniques, the effect of different types of somatic cells can be tested comparatively. In this way, more effective cells are identified during screening, after which the potent player can be identified. In that framework, the indirect effect of nutrients on cell development through somatic cells can be determined (65). Further, the bioactive components might be identified by serum or follicular fluid pre-fractionation (66, 67). For instance, follicular fluid has shown preserving effects on the stemness characteristics of human granulosa cells (68). In the approach of employing MSCs to produce PGCLCs, there is a great opportunity for manipulation before differentiation. So far, two studies (48, 52) have shown direct MSCs towards PGCLCs through genetic manipulation. With this strategy, the activity of related signaling pathways has been studied. Gene therapy can be performed once optimal PGCLCs are achieved. Thus, as with cases of reprogramming and transdifferentiation, genetically modified PGCLCs can be obtained from MSCs in genetic diseases.

At present, in vivo PGCLC grafting is mainly used to obtain gametes, as no suitable in vitro condition has yet been established for this purpose. For instance, Hayashi and Surani (63) successfully used somatic cells from mice embryonic oocytes to direct oogenesis from PGCLC, but the efficiency of oocytes derived from PGCLC to produce zygotes was only 53%. Providing appropriate conditions for the proliferation and maintenance of PGCLCs is a challenge in this regard. Many factors such as cell species, age, and sex affect MSC’s ability to eventually differentiate into gametes. Initial identification and purification of MSCs using exclusive and robust markers is an essential requirement for achieving optimal and defined conditions. Another important factor is the optimal duration of differentiation induction. Accordingly, it is necessary to determine the dynamics of markers in an integrated way. In addition, the appropriate stage of differentiation needs to be determined according to the ability of proliferation and subsequent differentiation into gamete lineages.

Conclusion

The availability of MSCs has made it possible to customize conditions for their differentiation into PGCLCs in several models, including humans. Umbilical cord, adipose tissue, and bone marrow are prospective sources of MSCs for germ cell line regeneration. Refining germ cell-related signaling pathways during induced differentiation of MSCs will help define extension to the protocols for PGCLCs production.

Acknowledgments

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Authors’ Contributions

Sh.F.; Contributed to the conception, design, and drafting. P.F., Z.N.; Contributed to the literature review. A.M.; Was responsible for overall supervision. All authors read and approved the final manuscript.

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The Effect of miR-106b-5p Expression in The Production of iPS-Like Cells from Mice SSCs during The Formation of Teratoma and The Three Embryonic Layers

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Abstract

Objective: According to the mounting data, microRNAs (miRNAs) may play a key role in reprogramming. miR-106b is considered as an enhancer in reprogramming efficiency. Based on induced pluripotent stem cells (iPSCs), cell treatments have a huge amount of potential. One of the main concerns about using iPSCs in therapeutic settings is the possibility of tumor formation. It is hypothesized that a procedure that can reprogram cells with less genetic manipulation reduces the possibility of tumorigenicty.

Materials and Methods: In this experimental study, miR-106b-5p transduced by pLV-miRNA vector into mice isolated spermatogonial stem cells (SSCs) to achieve iPS-like cells. Then the transduced cells were cultured in specific conditions to study the formation of three germ layers. The tumorigenicity of these iPS-like cells was investigated by transplantation into male BALB/C mice.

Results: We show that SSCs can be successfully reprogrammed into induced iPS-like cells by pLV-miRNA vector to transduce the hsa-mir-106b-5p into SSCs and generating osteogenic, neural and hepatoblast lineage cells in vitro as a result of pluripotency. Although these iPS-like cells are pluripotent, they cannot form palpable tumors in vivo.

Conclusion: These results demonstrate that infection of hsa-mir-106b-5p into SSCs can reprogram them into iPSCs and advanced germ cell lineages without tumorigenicity. Also, a novel approach for studying the generation of iPSCs and the application of iPSC or iPS-like cells in regenerative medicine is presented.

Keywords: Induced Pluripotent Stem Cells, Mir-106b, Spermatogonial Progenitor Cell, Transplantation, Tumorigenicity

Introduction

Yamanaka and colleagues were the first ones who produced adult fibroblasts into induced pluripotent stem cells (iPSCs) in a laboratory setting in 2006 providing the basis for substantial advancements in cell reprogramming technology (1, 2). Cell reprogramming has been used in developmental and stem cell biology and regenerative medicine fields for the last decade to investigate the potential of iPSCs in generating targeted cell types (3). In general, iPSCs may be produced in vitro from a variety of somatic cell types. Fibroblasts of mesodermal origins, endodermal hepatocytes, and ectoderm keratinocytes have been the most common cells used for this purpose till now (4-6). Spermatogonia stem cells (SSCs) are a type of testicular stem cell that has the ability to self-renew and differentiate into sperm cells. Current knowledge on biotechnology suggests that SSCs can be more efficient and safe in cell pluripotency studies over embryonic stem cell or adult somatic cell-based technologies. SSCs isolation from an individual’s testicular tissue eliminates ethical and immunological concerns in cell treatments. As reported in previous studies, generating iPSCs from adult fibroblasts is required retroviral transduction of pluripotent stem cell genes such Oct4, Sox2, klf4, c-Mys (OSKM), results in retroviral infection and teratomas. In contrast, SSCs can be reprogrammed to iPSCs in a specific culture medium without the addition of oncogenes or the use of retroviruses. However, tumorigenicity has been reported in this method (7, 8). According to new research, SSCs may be self-reprogrammed in a feeder-free reprogramming technique. Furthermore, SSCs can express the octamer-binding transcription factor4 (Oct4), a key factor in sustaining pluripotency in stem cells (9).

MicroRNAs (miRNAs) are a type of small non-coding RNA that are functional in the self-renewal, pluripotency, and differentiation of human embryonic stem cells (hESCs). MiRNAs play a crucial role in animal development by targeting mRNAs and regulating genes post-transcriptionally (10). Some miRNAs, including miR-106 increased reprogramming efficiency. miR-106b was reported one of the miRNAs with the highest expression of OSKM. These factors are known as the main factors that can be used for reprogramming (11). miR-106b belongs to the polycistronic miR-106b25
cluster and is found inside an intron of the MCM7 gene; it can significantly improve iPSC efficiency and boost the reprogramming process by targeting Tgfbr2 and p21 (12, 13).

According to reports from several laboratories, Yamanaka-induced pluripotency cells have the capacity to generate teratomas (14, 15). Probably, this specification is related to extensive genetic manipulation and the use of multiple viral vectors. Tumorigenicity has led to the limitation of the application of iPSCs in medicine, so a technique that can reprogram cells with less genetic manipulation reduces the possibility of tumorigenicity. According to the capacity of SSCs in converting to pluripotent cells and features of miRNAs mentioned above, we investigated the ability of reprogrammed SSCs into iPSC-like cells by pLV-miRNA vector to transduce the hsa-mir-106b-5p into SSCs and generating three germ layers (ectoderm, endoderm, and mesoderm). Furthermore, we studied the capability of these iPSC-like cells in tumorigenicity by measuring the size and pathology of tumors caused by the subcutaneous injection of cells into mice.

Material and Methods

Animals

Male BALB/C mice 8-10 weeks old and weighing 18-24 g, were treated with cyclosporine in a dose of 10 mg/kg per day by gavage. Transplanted mice received cyclosporine until two weeks after transplantation. Mice were maintained under sterile conditions on a 12-hour light-dark cycle at a constant temperature. Food and drink were freely available. All experiments were approved by the Ethics Committee of Shahid Beheshti University of Medical Sciences and following the Declaration of Helsinki (IR.SBMU.REC.1398.072).

Isolation and culture of SSCs

For the isolation of SSCs, testis of BALB/C mice suppressed by cyclosporine were collected and also isolation was performed by enzymatic digestion as described in our previous study. Spermatogonia cells were cultured for one week. According to the previous protocols explained in our earlier study, an antibody directed against promyelocytic leukemia zinc finger (PLZF) was used to identify the SSCs. Cells were identified with primary and secondary antibodies that were labeled with fluorescent reagents (16).

The pLV-miRNA vector production in bacteria

The pLV-miRNA vector, which carried the hsa-mir-106b lentivirus and comprised green fluorescent protein (GFP) in infected E. coli BL21, was utilized to generate iPSC-like cells (mir-p081, Biosettia, San Diego, CA, USA). The E. coli BL21 colony was cultured in 5 ml of Lysogeny broth (LB) medium (Sigma-Aldrich, USA) for 16 hours in a 37°C shaker incubator at 180 rpm. To determine the presence of vector in the bacteria, it was cultured for 24 hours in LB agar medium containing 100 µg/ml ampicillin. Vector purification from E. coli BL21 colonies was deployed by GF-1 Plasmid DNA Extraction Kit (Vivantis, Malaysia) instructions.

The quantitative reverse transcriptase polymerase chain reaction (qRT-PCR) technique was applied to ensure the accuracy of the extracted pLV-miRNA vector. The 16s-RNA as a bacterial reference gene and mir-106b primers are listed in Table 1. The sequences of forward and reverse primers were designed by using GeneRunner software. The cDNA was synthesized according to the manufacturer’s instructions (Fermentas, USA). PCR products were electrophoresed on 1% agarose gel along with 1 kb DNA Ladder Marker and examined.

Transduction of the hsa-mir-106b-5p vector to SSC colonies

To vector transduction, SSCs are inserted into the cells of a cell culture well. Then, in a microtube, 500 µl of culture media and 7.5 µl of lipofectamine 3000 (Invitrogen, USA) are combined. In a separate microtube, 500 µl of media is combined with 0.5 µg of plasmid and 7.5 µl of lipofectamine 3000. The contents of the two microtubes were combined and incubated at room temperature for 10 to 15 minutes. Finally, 250 µl of the prepared sample was pipetted into each plate housing, and the cells were incubated for three days at 37°C. It is important to validate the increase in miRNA after transduction of the virus carrying the hsa-miR-106b-5p gene into the cells. A fluorescent microscope was used to confirm viral transduction since the plasmid contains a GFP tracer reagent (Olympus BX51).

<table>
<thead>
<tr>
<th>Primer</th>
<th>Primer sequence (5’– 3’)</th>
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<tbody>
<tr>
<td>16s-RNA</td>
<td>F: ACTCTCACGGGAGGCAGCAG</td>
</tr>
<tr>
<td></td>
<td>R: ATTACCGCGGCTGCTGG</td>
</tr>
<tr>
<td>Stem loop</td>
<td>GGTGGCTCTGGTCAGGGTCCAGAGGTATTCGCACCAGTGCAANNNNNN</td>
</tr>
<tr>
<td>miR106b</td>
<td>F: ACUGCAGUGCCAGCAGCTT</td>
</tr>
<tr>
<td></td>
<td>R: GGCAAAAGTGCTTACAGTGCA</td>
</tr>
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Production of iPSCs Using miR-106b-5p

**Examination of iPSC-like cell differentiation into all three germ layers in vitro phase**

Differentiation into three germ layers was performed in two groups of cells: hsa-mir-106b-5p induction and without-hsa-mir-106b-5p induction as control group (treated with the empty vector). Both experimental groups were placed in a hanging drop culture, a concentration of $5 \times 10^4$ cells/ml suspension was prepared to soak in 20 µl drops to produce the embryoid body (EB). Procedure includes holding cells in a droplet of culture medium and turning the microplate upside down to produce 3D spheroids. Surface tension forces and gravity are two factors that keep cells suspended (17). At the end of the procedure in each experimental group, the differentiation of iPSC-like cells was observed in 5 randomly chosen fields under a fluorescence microscope.

**Differentiation into the ectodermal derivative**

Cells were transferred into a 12-cell gelatinized microplate containing α-minimum essential medium (α-MEM, Sigma-Aldrich, USA) with 3% fetal bovine serum (FBS, Gibco, UK). Toward induction of neural phenotype in the EBs in two weeks, the $5 \times 10^{-7}$ molar concentration of retinoic acid (Sigma-Aldrich, USA) was administered. Two weeks later, a beta-tubulin marker (Santa Cruz, USA) was used to examine the differentiation of adult neural cells (18).

**Differentiation into the mesodermal derivative**

Cells were grown in a six-cell plate when the entire culture surface was covered with cells, and their media changed with a bone differentiation medium to generate osteogenic lineage cells as a mesodermal lineage in transduced cells. This medium included Dulbecco’s Modified Eagle Medium (DMEM, Gibco, UK) with 10% bovine serum (Gibco, UK), 10 mM beta glycerol phosphate (Sigma-Aldrich, USA), 10 nM dexamethasone (Sigma-Aldrich, USA), and 50 g/ml ascorbic 3-phosphate (Sigma-Aldrich, USA). The cells were then put on mesenchymal cells by using a sampler. For 21 days, cells were cultured in a humidified 37°C incubator with 5% CO$_2$. Finally, immunocytochemistry was performed to validate the differentiation of cells using the alkaline phosphatase marker (Santa Cruz, USA) (19).

**Differentiation into the endodermal derivative**

Cells were cultured for 28 days in DMEM medium (Gibco, UK) containing 20 µl ascorbic acid (Sigma-Aldrich, USA) 20 µl, 10 ng/ml hepatocyte growth factor (HGF, Merck, Germany), 10 ng/ml oncostatin M (OSM, Sino Biological, China), 10% FBS (Gibco, UK). Immunocytochemistry was used to establish the presence of the albumin marker (Santa Cruz, USA) that enabled the cells to differentiate into hepatocytes (20).

**Tumorigenicity of cells**

Cells were transplanted into 20 immunodeficient male BALB/C mice (4-6 weeks). Transplantation was applied in four groups (n=5): SSCs (negative control), iPSCs as a positive control were obtained by using the method described by Baharvand and colleagues (21). They were provided by Stem Cells Technology Research Center. hsa-mir-106b-5p control (SSCs with empty vector), and hsa-mir-106b-5p (SSCs infected with hsa-mir-106b-5p). $5 \times 10^6$ µl of culture medium was transplanted by subcutaneous injections into the loose skin over the mice back, and assessed tumor formation after eight weeks. Generation of tumors were measured by a caliper. The tumors were then isolated and stained for pathological examination using the hematoxylin and eosin technique.

**Results**

**Confirmation of the nature of SSCs**

Cell culture experiments were performed with male BALB/c mice testis. SSCs were isolated from seminiferous tubules of the testis and cultivated in DMEM conditions. In the testis, PLZF is a spermatogonia-specific transcription factor that is detected to identify SSCs (22). Immunocytochemistry analysis demonstrated high purity and proliferation of SSCs by the expression of the PLZF marker (Fig.1A).

**Expression of mir106b in transfected bacteria**

The presence of the mir106b was determined by the qRT-PCR method. Results indicated a significant expression in mir106b in the transfected *E. coli* group compared to the non-transfected *E. coli* group.

**Confirmation of transduced cells**

Transduction of hsa-mir-106b-5p into SSCs confirmed by tracing the GFP protein reagent with a fluorescence microscope (Fig.1B).

**Differentiation of iPSC-like cells into all three germ layers**

Differentiation into the ectoderm derivative

The biomarker class III beta-tubulin is known for expresses in neural lineage cells (23). Therefore, a beta-tubulin marker was used to identify neurons in cultured cells. Immunocytochemistry analyzes revealed that the hsa-mir-106b-5p induction cell group differentiated into neurons, but there was no differentiation to neural cells in the without- hsa-mir-106b-5p induction cell group (Fig.2).
Fig. 1: Confirmation of SSC nature as well as transfect. A. Immunocytochemical characterization of promyelocytic leukemia zinc finger (PLZF) in cultured spermatogonia stem cells (SSCs). Immunocytochemistry analysis of PLZF expression in the SSC (scale bar: 50 µm). B. Tracing the GFP protein reagent with a fluorescence microscope to confirm transduction. The phase contrast, fluorescent, and merged photomicrographs demonstrate the transduction of hsa-mir-106b-5p into SSCs (scale bar: 40 µm).

Differentiation into the mesoderm derivative

As shown by immunocytochemistry analyzes of alkaline phosphatase (ALP) in Figure 3, iPS-like cells with induction of hsa-mir-106b-5p were differentiated to osteogenic lineage cells. ALP is a frequently used marker for observing cell lineages of the embryonic mesoderm such as osteogenic lineage (24). However, osteocyte differentiated cells were not detected in the group without hsa-mir-106b-5p induction.

Fig. 2: Fluorescent immunocytochemistry of beta-tubulin marker to detect ectodermal derivatives. hsa-mir-106b-5p induction group: differentiated neuronal cells exhibited. Without-hsa-mir-106b-5p induction group: no differentiation detected (scale bar: 20 µm).

Differentiation into the endoderm derivative

Adult functional hepatocytes secrete albumin into the culture medium which is used as a marker to identify endoderm lineage cells (25). Immunocytochemistry analyzes in Figure 4 showed that at the completion of the procedure, in the hsa-mir-106b-5p induction cell group, hepatocytes produced intracellular albumin but were incapable to released albumin into the extracellular environment. As a result, the cells were hepatocyte-like and developed into hepatocyte lines. However, the differentiated cells were not functional.

Fig. 3: Fluorescent immunocytochemistry of alkaline phosphatase marker to detect mesodermal derivatives. hsa-mir-106b-5p induction group: differentiated osteocyte cells exhibited. Without-hsa-mir-106b-5p induction group: no differentiation detected (scale bar: 20 µm).

Fig. 4: Fluorescent immunocytochemistry of Albumin marker to detect endodermal derivatives. hsa-mir-106b-5p induction group: Albumin marker expressed in the hepatocyte-like cells but not detected in the secretion of the cells. Without-hsa-mir-106b-5p induction group: no differentiation detected (scale bar: 20 µm).

Tumor growth in mice

Four independent experiments were performed, each using a different colony of have transduced SSCs and none-transduced cells. In three experimental groups [the SSCs negative control], has-mir-106b-5p control (empty vector), and has-mir-106b-5p (SSCs infected with hasa-mir-106b-5p) no palpable tumors were observed at the end of the experiment after two months. In contrast, the
iPSCs (positive control) formed palpable tumors at the site of injection in the mice back. Histological examination of tissue sections revealed tumorigenicity that accrued by iPSCS injection and the tumors grew 0.5×1×1 cm at week eight, when the animals were sacrificed (Fig.5).

Discussion

Studies have shown that induced pluripotency SSCs can differentiate in vitro into the three germ layers including endoderm, mesoderm, and ectoderm. SSCs can conveniently be induced into pluripotent stem cells in a specified culture medium (26, 27). Shinohara and colleagues reported in 2004 that they had generated embryonic stem-like cells (ES-like) from mouse testis SSCs that were phenotypically resembling to ES cells (28). Reprogramming competence of SSCs qualified them into the proper cells for iPSCs in rodents and human research (29). These iPSC cells produce teratomas after transplantation which is the most significant obstacle of using these cells in medicine. Our findings demonstrate that induction of pluripotency in SSCs can be achieved by inducing hsa-mir106b-5p. These iPSC-like cells could establish embryonic lineages. Also, these cells did not show tumorigenicity, which is their privilege compared to other reprogrammed cells.

Tumor creation in iPSCs is related to the activation of some oncogenes such as c-Myc. Nakagawa et al. (30) reported that removing c-Myc from the reprogramming process resulted in the development of pluripotent cells but eliminated teratoma formation. We have observed SSCs can be successfully reprogrammed into iPSC-like cells using procedures without multiple retroviral transductions. It can be concluded that hsa-mir-106b-5p does not target c-Myc gene, and we generated pluripotent cells without tumorigenic properties. Previously, it was reported that the miR-106b-5p cluster significantly increased during reprogramming stages, inhibiting this cluster decreases reprogramming efficiency, so it is a reprogramming activity regulator (12). In addition, studies showed that certain miRNA families, such as miR-17, miR-106, miR-520, miR-372, miR-195, and miR-200, are upregulated in human pluripotent stem cells (hPSCs) as compared to adult differentiated cell types. Moreover, miRNA clusters can promote reprogramming into iPSCs. The miR-106b-25 cluster is proven to be early active in the reprogramming of mouse embryonic fibroblasts (31-34). Lin and colleagues induced mir-302 into human hair follicle cells by an inducible vector; this procedure resulted in some human embryonic specific gene expression, such as NANOG, OCT3/4, SOX2 so these somatic cells were successfully reprogrammed to iPSCs (35). Nguyen and colleagues revealed that co-expression of miR-524-5p with OSKM factors in human somatic cells leads to generate iPSCs. According to their study, miR-524-5p initiates reprogramming by targeting ZEB2 and SMAD4 genes, which are epithelial-to-mesenchymal transition-related genes (36). Based on the functional role of mir-106b in cell reprogramming, the dosage of hsa-mir-106b, a naturally present miRNA in spermatogonial stem cells was enhanced in this technique by infecting the pLV-miRNA vector into isolated SSCs to generate iPSCs (37). Here, these iPS-like cells showed pluripotent characteristics and successfully differentiated into osteogenic cells (mesoderm derivative), neuronal cells (ectoderm derivative) and hepatocyte lineage (endoderm derivative) as a result of reprogramming. Isolated SSCs from adult mice have been shown to be capable of reprogramming and differentiating into all three embryonic germ layers under three defined culture conditions, as well as generating teratomas in immunodeficient mice (38). Another study by Lim and colleagues reported in vitro expression of three germline markers in the EB-like structures in iPSCs of human SSCs (39). These findings authenticate that induction of pluripotency in SSCs can lead them to produce embryonic layers. Although pluripotency of iP cells is a remarkable result in cell-based therapies, the tumorigenicity of this type of cells is still a concern for clinical applications (40). Based on findings in this study, transduction of hsa-mir106b-5p to SSCs by one vector and limited genetic manipulation led to reprogramming of SSCs into pluripotent cells without the capability of tumor formation. Through this method, there can be a new prospect at the generation of iPSCs and the application of iP or iP-like cells in regenerative medicine. This method can be an acceptable alternative.
for techniques that are involved with substantial genetic manipulation. Besides, vectors as a viral basis element are not allowed to use considerably in the human body, so we can take advantage of them in a minimum of manipulation to produce pluripotent cells without tumorigenicity.

Conclusion

The results of this study showed that using this new method in infecting hsa-mir-106b-5p into SSCs, leads to reprogramming them and turning them into pluripotent cells. iPS-like cells differentiated successfully into germ layers as a result of pluripotency. On the other hand, iPSC-like cells do not cause tumors, which is a significant characteristic of iPSCs in medical applications. This reprogramming method provides a simplified and convenient way to convert SSCs into pluripotent cells with less ethical and immunological concerns in cell treatments.

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Authors’ Contributions

A.H.F., Z.M., S.J.H.; Proposed and performed experimental works, and data collection. F.K.; Performed bioinformatics work. F.K., A.H.F.; Contributed to article writing and manuscript approving. All authors read and approved the final manuscript.

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Production of iPSCs Using miR-106b-5p


Differentiation of Alginate-Encapsulated Wharton Jelly-Derived Mesenchymal Stem Cells into Insulin Producing Cells

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Abstract

Objective: Insulin insufficiency due to the reduced pancreatic beta cell number is the hallmark of diabetes, resulting in an intense focus on the development of beta-cell replacement options. One approach to overcome the problem is to search for alternative sources to induce insulin-producing cells (IPCs), the advent of mesenchymal stem cells (MSCs) holds great promise for producing ample IPCs. Encapsulate the MSCs with alginate improved anti-inflammatory effects of MSC treatment. This study aimed to evaluate the differentiation of wharton jelly-derived MSCs into insulin producing cells using alginate encapsulation.

Materials and Methods: In this experimental study, we established an efficient IPCs differentiation strategy of human MSCs derived from the umbilical cord’s Wharton jelly with lentiviral transduction of Pancreas/duodenum homeobox protein 1 (PDX1) in a 21-day period using alginate encapsulation by poly-L-lysine (PLL) and poly-L-ornithine (PLO) outer layer. During differentiation, the expression level of PDX1 and secretion of insulin proteins were increased.

Results: Results showed that during time, the cell viability remained high at 87% at day 7. After 21 days, the differentiated beta-like cells in microcapsules were morphologically similar to primary beta cells. Evaluation of the expression of PDX1 and INS by quantitative reverse transcriptase-polymerase chain reaction (qRT-PCR) on days 7, 14 and 21 of differentiation exhibited the highest expression on day 14. At the protein level, the expression of these two pancreatic markers was observed after PDX1 transduction. Results showed that the intracellular and extracellular insulin levels in the cells receiving PDX1 is higher than the control group. The current data showed that encapsulation with alginate by PLL and PLO outer layer permitted to increase the microcapsules’ beta cell differentiation.

Conclusion: Encapsulate the transduced-MSCs with alginate can be applied in an in vivo model in order to do the further analysis.

Keywords: Alginate, Diabetes, Insulin, Mesenchymal Stem Cells


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Introduction

Diabetes is one of the metabolic disorders which are caused by impaired insulin secretion, insulin dysfunction, or both. Numerous pathogenic processes contribute to the progression of diabetes, which has a wide range from progressive autoimmune destruction of pancreatic cells toward ultimately insulin deficiency and insulin resistance. Until 2017, 387 millions of people worldwide suffer from diabetes, and that number is expected to reach in to 500 millions by 2030 (1). These statistics have led the international diabetes federation (IDF) in to describe this disease as one of the most serious human health challenges in the 21st century (2).

So far, no certain cure option for diabetes currently exists. At present, taking oral medications and insulin injections are common treatments. Transplantation of islets isolated from the donor pancreas could also be a therapy for diabetes. However, this treatment also has some restrictions, including limited islets required for transplantation, side effects of long-term immunosuppressive drugs, and short-term survival after transplantation. Recently, the application of stem cells that differentiate into pancreatic beta cells has drawn great attention as one of the treatment options for diabetes (3). Some studies suggested that embryonic stem cells, induced pluripotent stem cells (iPSCs), bone marrow-mesenchymal stem cells (BM-MSCs) and adipose tissue-MSCs can differentiate into IPCs both in vitro and in vivo (4). Stem cells with the ability to differentiate into insulin-producing cells (IPCs) are becoming the most promising therapy for diabetes mellitus that reduce the major limitations of availability and allogeneic rejection of beta cell transplantations (5, 6).

Previous studies have reported that iPSCs were derived from embryonic stem cells in mice and humans (7, 8). MSCs have potentials for differentiating into various tissues, immunomodulatory effects and the invasiveness of the procedure. Therefore MSCs can overcome the obstacles seen with embryonic stem cells (9).

MSCs are the most important candidates for cell
therapy which are obtained from different sources including BM, adipose tissue, blood, amniotic fluid and umbilical cord of newborns. Recently umbilical cord derived MSCs from Wharton jelly have drawn many attentions because of their differentiation, migration and protective properties compare to other kinds of stem cells. Hu et al. (10) evaluated the application of umbilical cord’s Wharton jelly-derived MSCs (WJ-MSCs) for type 1 diabetic patients and obtained promising results. Another study established that transplantation of placenta-derived MSCs for patients with type 2 diabetes was safe, easy, and potentially effective (11).

Alginate hydrogels have demonstrated high applicability as a structure for cell immobilization. Alginate is recognized in properties such as its ability to make hydrogels at physiological conditions, gentle dissolution of gels for cell retrieval, transparency for microscopic evaluation, gel pore network that allows diffusion of nutrients and wastes in addition to its reduced risk of graft failure (12). Encapsulation is a method used to protect implanted cells from immune system attack and it may enhance the survival rate and differentiation of implanted cells by the increased of cytokines secreted by encapsulated cells to the microenvironment (6, 13). Microencapsulation is widely used for encapsulation of cells or bioactive molecules, gene therapy and drug delivery. Hydrogels are the most widely used materials for cell microencapsulation because of their high porosity that leads to high permeability of oxygen, nutrients, and metabolites (14). Alginate widely used for cell encapsulation provides protection of the encapsulated cells against the host’s immune system (15). Previous studies demonstrated that MSC encapsulated in alginate could survive locally after implantation in vivo (16).

The transplantation of pancreatic islets in immune protective capsules holds the promise as a functional cure for type 1 diabetes (17), about 40 years after the first proof of principal study (18). Gene therapy, as an advanced technology to treat diseases cannot be treated with conventional medicine and can be applied to a wide range of diseases that includes many methods of gene transfer (19). Gene therapy had been approved for diseases such as cystic fibrosis, diabetes, autoimmune diseases, heart diseases Alzheimer’s disease, Parkinson’s disease, various cancers (20). Gene therapy by viral vector and non-viral transduction may be useful techniques to treat diabetes (21). Insulin generation in MSCs through genetic engineering is a promising therapeutic for patients with diabetes (22). In previous study it was indicated that PDX1-tranduced hBM-MSCs differentiate into IPCs (21). This study aimed to evaluate the differentiation of wharton jelly-derived MSCs into insulin producing cells using alginate encapsulation.

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<th>Material and Methods</th>
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<td><strong>Isolation of MSCs from Wharton jelly</strong></td>
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<td>In this experimental study, umbilical cords were collected from healthy full-term deliveries after receiving consent from parents. The collection and using of human biological specimens were approved by the Ethics Committee of the Islamic Azad University-Science and Research Branch (IR. IAU.SRB.REC.1398.214). The Umbilical cords were transferred in serum-free Dulbecco’s Modified Eagle Medium/F12 (DMEM/F-12, Hyclone, Logan, UT, USA) and transferred to the laboratory immediately. After washing, the Umbilical cords samples were cut into 2-3 cm sections, the umbilical vessels removed, and Wharton jelly was collected and minced into pieces. The pieces were plated in tissue culture flasks containing an enzymatic solution of collagenase and hyaluronidase, in DMEM/F-12 medium supplemented with 10% fetal bovine serum (FBS, Gibco, USA) and incubated at 37°C in a humidified 5% CO₂ incubator for 45 minutes to 2 hours. This allows Wharton jelly loosening and separation from the Umbilical cords without complete digestion. After the incubation period, the Umbilical cords pieces are transferred to a new Petri dish or culture flask containing fresh DMEM to remove any remaining enzymes (23).</td>
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<th>Material and Methods</th>
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<td><strong>Flow cytometry analysis</strong></td>
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<td>Human MSCs single-cell suspensions were harvested using a 0.05% trypsin/ Ethylenediaminetetraacetic acid (EDTA) solution; after FBS neutralization incubated in blocking buffer [1% FBS in Dulbecco’s phosphate-buffered saline (DPBS)] for 30 minutes. Next, 1×10⁶ cells were separately incubated for 1 hour at 4°C with an optimal dilution of conjugated antibodies that included anti-CD73-FITC (ab28061), anti-CD45-FITC (ab27287), anti-CD90-FITC (ab11155), anti-CD34-PE (ab157304), and anti-CD105-PE (ab91138), all from Abcam (Cambridge, UK). Flow cytometry experiments were performed with a BD FACSCalibur Flow Cytometer (BD Biosciences) and data analyzed by the Flowing software.</td>
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<td><strong>Multilineage differentiation</strong></td>
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<td>To confirm the multipotency of WJ-MSCs, osteogenic and adipogenic differentiation were verified with alizarin red and oil red O staining respectively. To induce osteogenesis, WJ-MSCs treated with osteogenic differentiation medium, alpha minimum essential medium (Life Technologies, USA) supplemented with 10% FBS (Gibco, USA), 10 mmol/L β-glycerophosphate (Sigma-Aldrich, USA), 0.1 mmol/L dexamethasone (Sigma-Aldrich, USA) and 50 mmol/L ascorbic acid (Sigma-Aldrich, USA) for 21 day. For adipogenesis, the cells were treated with adipogenic differentiation medium</td>
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in alpha minimum essential medium supplemented with 10% FBS, 1 mmol/L dexamethasone (Sigma-Aldrich, USA), 10 mg/mL insulin (Sigma-Aldrich, USA), 0.5 mmol/L isobutyl-methylxanthine (Sigma-Aldrich, USA) and 100 mmol/L indomethacin (Sigma-Aldrich, USA) for 21 day.

Transduction of WJ-MSCs

To transduce PDX1 using lentivirus system, approximately 1×10⁶ MSCs were seeded in 48-well plates. The Lentivector Packaging kit (Invitrogen, USA, K4975-00) including the pPackH1 Packaging Plasmid (mixture of pPACKH1-GAG, pPACKH1-REV, and pVSV-G plasmids, 0.5 μg/μl) and the transfer vector Plenti-Pdx1-PURO (0.5 μg/μl) containing an enhanced PDX1, was used to transduce PDX1 into WJ-MSCs. Virion particles were produced in 293T cells (Invitrogen, Carlsbad, CA, USA) by transfection using the TransIT-2020 Transfection Reagent (Mirus, Madison, WI, USA). The 293T cells were seeded in 75-cm² flasks at an initial density of 1.3×10⁶ cells/cm² with 10 ml of DMEM containing 10% FBS, 50 U/ml penicillin, and 50 μg/ml streptomycin. At 24 hours post-transfection, the media was replaced with fresh DMEM with 2% FBS. The medium was changed every 24 hours for 3 days. The media was removed, pooled, and filtered (pore size: 0.45 μm; Merck Millipore, Rockland, UK), 20 ng/ml β-fibroblast growth factor (bFGF, Sigma-Aldrich, USA), 2% B27 (Gibico, UK), 2 mmol/L L-glutamine (Hyclone, USA), 10 ng/ml β-cellulin (Sigma-Aldrich, USA), 10 ng/ml activin A (Sigma-Aldrich, USA), 2% B27 and 10 mmol/L nicotinamide (Sigma-Aldrich, USA) (7) were used as negative control, and PANC-1 cell line (pancreatic epithelial cells) was used as a positive control (24).

Total cellular RNA was extracted by the TRIzol reagent® (Sigma-Aldrich, T9424) and used for cDNA synthesis with the Revert Aid First Strand cDNA Synthesis Kit (Fermentas, Germany, K1632) according to the manufacturer’s instructions. Quantitative RT-PCR was carried out with the SYBR Green Master Mix (Takara Bio, Inc., RR081Q) with a real-time RT-PCR system (Corbett Life Science, Rotor-Gene 6000, Australia). The expression levels of the target genes were calculated using the 2⁻ΔΔCt method with glyceraldehyde 3-phosphate dehydrogenase (GAPDH) as the internal control for normalization. Primer sequences for target genes are listed in Table 1.

Western blotting

For western blot analysis at the end of treatment, on day 21, the cells were lysed in commercial lysis buffer (Qproteome Mammalian Protein Prep Kit, QIAGEN) according to the manufacturer’s protocol. The solubilized protein fractions of each sample (50 μg) from three biological replicates were separated on a 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and transferred onto a PVDF membrane (Amersham Biosciences, USA) by semi-dry blotting (Bio-Rad, USA) using transfer buffer (10 mM NaCHO₂, 3 mM Na₂CO₃, 20% methanol). Membranes were blocked with Tris-buffered saline with Tween® 20 (TBST, 20 mM tris-HCl, pH=7.6, 150 mM NaCl, and 0.1% Tween-20) that contained 5% BSA and then incubated overnight with the primary antibody at 4°C. After three times washing with TBST, the membranes were incubated with horseradish peroxidase (HRP)-conjugated secondary antibody at room temperature for 1 hour. Signals were detected with ECL substrate using Hyperfilm. Protein band intensity was normalized to the level of beta-actin. Each experiment was repeated at least three times.

Encapsulation of WJ-MSCs

The encapsulation of WJ-MSCs was performed as reported by Kanafi et al. (25), with slight modifications. An alginate solution (2% w/v) was prepared by dissolving 2 g of low viscosity alginic acid sodium salt (Low viscosity 100-300 cP, Sigma-Aldrich, USA) in 100 ml of deionized water. The alginate solution was mixed by overnight vortexing. To prepare cell-encapsulated beads, WJ-MSCs at passage 4 were harvested and a cell density of 5×10⁶ was mixed in 1 ml of alginate solution. The alginate solution was transferred to a 5 ml syringe (22 Gauge) and then extruded dropwise into an ice-cold 100 mM calcium
chloride (CaCl₂, Sigma-Aldrich, USA) solution. The droplets were left 10 minutes in the CaCl₂ solution for polymerization. Cell-encapsulated microcapsules were then transferred to 35 mm tissue culture dishes containing 1.5 ml DMEM medium supplemented with FBS. The WJ-MSC beads were incubated at 37°C in 5% CO₂ incubator for 72 hours and then used for further experiments.

### Decapsulation

Three days after encapsulation, microcapsules containing WJ-MSCs were washed by PBS twice, and 10 ml of decapsulation solution (EDTA, 50 mM and HEPES, 10 mM in PBS, Sigma-Aldrich, USA) was added, then the beads were incubated at 37°C for 10 minutes. The cells were pelleted by centrifugation at 3000 rpm for 10 minutes. This cell pellet was used for RNA isolation or protein extraction.

### Cell viability assessment

Assessment of WJ-MSCs viability was performed using the MTT assay. The cells at a density of 2×10⁵ well were inoculated into a 96-well plate. The WJ-MSCs were divided into transduce and transduced groups. The plates were placed in an incubator at 5% CO₂ at 37°C overnight. WJ-MSCs were transduced with Plenti-Pdx1-PURO lentivirus (5×10⁹ TU/ml) at 200 MOI based on the results of transfection efficiency, and WJ-MSCs in the untransfected group received an equivalent dose of PBS. Cells in one of the four plates were incubated with the MTT solution at 7, 14, and 21 days after transfection. After 4 hours, the medium was removed, and 150 μl of dimethylsulfoxide (DMSO) added to each well. Absorbance was measured at 570 nm using an ELISA reader (Biochrom Anths 2020, UK).

### Insulin assay and response to glucose

Insulin level in the medium was measured by human insulin ELISA kit (Millipore, Billerica, MA, USA) according to the manufacturer’s instructions. Total protein in the medium was measured by the BCA assay using fresh culture medium as a blank. To determine the cell response to glucose at different concentrations, the insulin levels were evaluated with different glucose concentrations (0, 5.5, 15 and 25 mM). 1×10⁶ cells were initially incubated for 3 hours in glucose-free Krebs-Ringer bicarbonate buffer (KRB). This was followed by incubation for 1 hour in 3.0 mL of KRB containing 0, 5.5, 15, or 25 mM glucose concentrations. The supernatant was collected at the end of each incubation period. The collected samples were using the ELISA assay (26). We evaluated the insulin level in positive control group to confirm the insulin ELISA kit.

### Statistical analysis

All experiments were conducted in at least three independent repeats and performed in the same passage. Statistical analysis was performed using GraphPad Prism 5.02 (GraphPad Software, Inc, USA). Comparisons between groups were performed by one-way analysis of variance (ANOVA) followed by the Tukey post-hoc test. The independent t test analysis was carried out to identify statistical differences between the two observations. The difference between data was considered to be significant at P<0.05.

### Results

#### Derivation and characterization of WJ-MSCs

MSCs derived from human umbilical cord’s Wharton Jelly, human WJ-MSCs had fibroblastic-like phenotype (Fig.1A), the cells were small and fusiform at the first passage. After third passage, the cells seem fully expanded with many cytoplasmic processes. To confirm the mesenchymal identity, the expression of MSC-specific markers was examined. Results from flow cytometry showed that the expression (%) of these markers including CD105, CD90 and CD73 in WJ-MSCs were 97.4, 96.70 and 95.3, respectively. While hematopoietic specific markers such as CD34 and CD45 did not have significant expression in these cell populations. These results confirm that the isolated cells from human umbilical cord’s Wharton jelly are MSCs (Fig.1B). WJ-MSCs are determined as multipotent stem cells that are able to differentiate into specific lineages like osteoblastic and adipocytic. Thus, the osteogenic differentiation assay was performed to examine the differentiation ability of isolated MSCs into these two lineages. Intracellular lipid droplets staining using oil red-O showed the adipogenesis of WJ-MSCs. While in the undifferentiated WJ-MSCs, these observations were absent. Alizarin staining demonstrated the formation of calcium oxalates on the differentiated MSCs, which was not detected in the undifferentiated cells. These findings confirmed the characterization of cells as WJ-MSCs and indicate that the MSCs have potential to differentiate into these lineages (Fig.1C).

#### Differentiation of MSCs derived from Wharton jelly into IPCs

To investigate whether transduction of WJ-MSCs with PDX1 leads to their differentiation into IPCs in vitro, we used the lentiviral vector to transfer the PDX1 gene into MSCs (Fig.2A). For this, WJ-MSCs were cultured in 6-well plates at 1×10⁶ cells/well (at passage number 3), when the confluency were reached 70-80%, transduction was performed (Fig.2B). After selecting the transduced cells using puromycin (at a concentration of 2 mg/ml),
these cells were cultured in serum-free newly culture medium for 21 days (Fig.2C). To evaluate the transduction efficiency of target cells, we evaluated the expression of beta cell-specific genes \( PDX1 \) and \( INS \) by qRT-PCR on days 7, 14 and 21 of culture (Fig.3A-C). We found that the expression levels of \( PDX1 \) and insulin genes in the PDX1-transduced group were higher than the negative control groups. Whereas, the difference between PDX1-transduced group and positive control groups (pancreatic cell) was not significant. On day 7, \( PDX1 \) and \( INS \) showed the lowest expression, and on day 14, exhibited the highest expression. We also examined the expression of these genes at the protein level by Western blotting; the expression of these proteins was examined 21 days after \( PDX1 \) transduction of WJ-MSCs. The results showed that PDX1 protein level was increased in transduced group (Fig.3D).

Fig.1: Isolation of MSCs from Wharton jelly, culture, and identification. A. Culture of WJ-MSCs during 21 days (scale bar: 100 µm). B. Evaluation of CD markers by flow cytometry. The WJ-MSCs expressed CD105, CD90, and CD73 but they expressed CD34 and CD45 at very low level. Each cell treatment was assayed on three technical replicates on three different samples of WJ-MSCs. C. Alizarin red staining after 21 day of culture in osteogenic medium indicated the osteogenic differentiation potential of WJ-MSCs (scale bar: 50 µm). Oil red staining after 21 day of culture in adipogenic medium showed the adipogenic differentiation potential of WJ-MSCs. Data for each day represent mean cells number and error bars show standard error of the mean (SEM) of triplicate experiment (n=3). MSC; Mesenchymal stem cells and CD; Cluster of differentiation.

Fig.2: Differentiation of WJ-MSCs into IPCs. A. Schematic summary of transduction pdx1-planti in cultured WJ-MSCs. B. Morphology of WJ-MSCs were transduced with the PDX1 gene using the Lentiviral vector system (scale bar: 200 µm). WJ-MSCs; Wharton jelly-derived mesenchymal stem cells and IPCs; Induce insulin-producing cells.
WJ-MSC Can Differentiate into IPCs using Alginate Encapsulation

Viability, and intra- and extracellular insulin levels in encapsulated PDX1-tranduced MSCs

In order to evaluate the effect of cell encapsulation by alginate hydrogel, the cell viability was evaluated by MTT assay on days 7, 14 and 21 after encapsulation (Fig.4A, B). The results showed 98% viability in the group that the cells were decapsulated immediately after encapsulation. After 7 days of cell encapsulation, cell viability remained high at about 87%. However, this rate was decreased to 79% following 14 days due to repeated passages. The viability of transduced-MSCs significantly increased following encapsulation by alginate on day 21. At the following examinations, to evaluate the functionality of encapsulated differentiated cells carrying PDX1, intra- and extracellular insulin levels were measured on day 14 using ELISA assay. Results showed that the intracellular and extracellular insulin levels in the MSCs receiving PDX1 is higher than the control group at concentration of 5.5, 15 and 25 ng/mg insulin protein (Fig.5A, B).

Fig.3: Investigation of differentiation of WJ-MSCs into IPCs. A., B. C. Expression of pancreatic-specific genes PDX1 and INS was analyzed by qRT-PCR in differentiated cells on days 7, 14 and 21. MSCs that were not cultured in the differentiating medium were used as negative control, and PANC-1 cell line was used as positive control. Glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was considered as the housekeeping control. Each experiment was conducted with in triplicate (n=3). D. Western blot analysis for PDX1 and its phosphorylated form (pPDX1) in MSC-Pdx1. Values represent mean and error bars show standard error of the mean (SEM) of triplicate experiment (n=3). ***; P<0.001, WJ-MSCs; Wharton jelly-derived mesenchymal stem cells, IPCs; Induce insulin-producing cells, and qRT-PCR; Quantitative reverse transcriptase–polymerase chain reaction.

Fig.4: Encapsulated MSCs in alginate hydrogels. A. The microscopic image of the capsules containing MSCs showed the uniform distribution of the cells within hydrogel. The average diameter of the capsules is 650 μm (scale bars: 50 μm). B. Percentage of viable cells evaluated by MTT assay, after 7, 14 and 21 days (n=3). MSCs; Mesenchymal stem cells, NC; Negative control, *; P<0.05, **; P<0.01, and ***; P<0.001.
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Fig. 5: Insulin assay and response to glucose. A. The expression of level insulin protein in the differentiated cells compared to the undifferentiated cells. Conversion factor for insulin: 1 μg=23 mU/L, 1 mU=0.6 pmol/L. B. Concentrations of intracellular insulin protein content in differentiated cells compared to undifferentiated cells. Conversion factor for insulin: 1 μg=23 mU/L, 1 mU=0.6 pmol/L (n=3). Control group; Cells without encapsulation and transduction. ***: Significant.

Discussion

The prevalence of diabetes is steadily increasing worldwide, making it one of the most challenging health issues of the 21st century (2). Therefore, it is necessary to search new effective treatment strategies aimed at recovering lost IPCs and inhibiting autoimmune destruction of endocrine progenitor cells (27). Also, results of a meta-analysis by El-Badawy and El-Badri showed that MSCs could be beneficial in patients with type 1 and type 2 diabetes (28). Differentiation of BM-MSCs and adipose tissue-MSCs into IPCs provides a new and promising strategy to reconstitute pancreatic endocrine function (26). In line with previous studies, our results showed, WJ-MSCs were differentiated into IPCs. Therefore, WJ-MSCs are a promising source for applying in diabetes due to their availability, low cost, and immune-modulatory properties. To produce IPCs from WJ-MSC origin, two methods of indirect and direct differentiation are used (29, 30). Indirect differentiation is performed using chemicals (e.g., nicotinamide and growth factors) and direct differentiation is based on genetic manipulation (31). We showed that transfer of PDX1 gene with lentiviral vector caused the differentiation of WJ-MSCs into IPCs, and confirming obtained results, an increase in the expression of pancreatic-specific genes such as PDX1 and insulin was observed. The results of our study were comparable to the findings of Rahmati et al. (32), in which the transfer of PDX1 gene to mouse MSCs with lentiviral method led to the differentiation of these cells to IPCs. In a study by Soltanian et al. (33), assembling three-dimensional (3D) pancreatic organoids (containing human embryonic stem cell-derived PDX1-positive pancreatic progenitors, MSCs, and endothelial cells) implanted into the peritoneal cavity of immunodeficient mice where it remained for 90 days. Their results indicated that 3D organoids developed more vascularization and a higher number of insulin-positive cells and improvement of human C-peptide secretions. Lima et al. (34) through overexpression of three beta cell-specific genes PDX1, Neurogenin-3 (NEUROG3), and V-maf musculoaponeurotic fibrosarcoma oncogene homolog A (MAF4) by adenoviral vectors, transdifferentiated pancreatic exocrine cells into beta-like cells both morphologically and functionally. In this study, we indicated that PDX1-tranduced MSCs group differentiate into IPCs.

Besides, for a durable treatment of insulin-dependent diabetes mellitus, it is crucial to establish a functional system that, in addition to supporting the insulin secretion in response to different levels of glucose, protecting from immune system. The viability and function of differentiated WJ-MSCs is the most important issue in the use of these cells in diabetes therapy. In this study, alginate hydrogel was used for encapsulation of the cells to avoid a declined survival. One important characteristic of alginate is its very limited inherent cell adhesion and cellular interaction, that is an advantage for cell encapsulation applications (12). The technique to cell immobilization, particularly pancreatic islet cells, in calcium alginate matrices was developed by Lim and Sun (18). By coating the alginate gel bead with polycations like PLL, PLO, or chitosan, the strength of the surface coating as well as the capsule porosity can be controlled (12). In this study, for production of alginate droplets (microcapsules), the viscous solution of alginate was mixed with the cells and then these were stabilized by treatment with polycationic polymers. A combination of PLL and PLO was applied as polycation in order to improve the strength of outer surface of microcapsules. The essential requirements for cell culture i.e. porosity, stability and permeability were reached by alginate in MSCs differentiation into IPCs (35). Consistent results by some studies were reported that pancreatic islet, ESC- and iPSC-derived IPCs encapsulated using alginate could maintain the viability both in vivo and in vitro (36).
The differentiated WJ-MSCs which were encapsulated by alginate, showed an increased cell viability, however this value was decreased to 79% after 14 days. Thickening of the outer layer by polycationic polymers may also cause the insufficient nutrient and oxygen consumption, which results in a significant reduction in cell viability on day 21. However, despite declined cell viability, insulin secretion levels did not decrease on days 7, 14 and 21. The high number of viable cells in viscous alginate capsules can also lead to high insulin secretion. EncapsulationMSCs levels did not decrease on days 7, 14 and 21. However, despite declined cell viability, insulin secretion 

Similar results have been obtained from other studies. In a recent study, alginate solution improved survival and maintenance of cell functionality in encapsulation of BRIN-BD11 beta cell line, and also the expression of INS was increased by 66% (38). Recently, Kuncorojakti et al. (3) evaluated the encapsulation of human dental pulp-derived stem cells (hDPSCs)-derived IPCs by alginate and pluronic F127-coated alginate. Obtained results showed that alginate and alginate combined with pluronic F127 preserved hDPSCs viability and allowed glucose and insulin diffusion in and out. In hDPSC-derived IPCs maintained viability for at least 14 days and sustained pancreatic endoderm marker (NGN3), NKX6.1, MAFA, ISL-1, GLUT2 as pancreatic islet markers, and intracellular pro-INS and INS expressions for a 14-days period. In another study, differentiation of WJ-MSCs into IPCs using a lentiviral system containing the GFP reporter gene and its transmission to diabetic NOD mice showed an elevated level of serum insulin and an improved glucose tolerance. Mice treated with WJ-MSCs-GFP had significantly lower blood sugar and higher survival rates than control mice (39). Results from a recent study by De Mesmaeker et al. (40) showed that encapsulation of porcine islet cells by alginate hydrogel in microsphere form enabled long-term glycemic control in immune-compromised mouse model of diabetes. This intracapsular functional beta cell mass formation involved beta cell replication, significant increasing number, and maturation toward human adult beta cells.

Conclusion

Our results showed that the differentiation of WJ-MSCs into Insulin Producing Cells is increased in PDX1-tranduced MSCs group. The INS level in encapsulated PDX1-tranduced MSCs with alginate was increased compared to the control group. Therefore due to the ability of WJ-MSC in amelioration fibrosis, modulation inflammation and enhancement vascular growth, MSCs could offer a promising treatment option for patients with endocrine disorders.

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Authors’ Contributions

Z.P.; Participated in study design, data collection and evaluation, drafting and statistical analysis. S.K., N.H.R.; Were involved in data analysis and interpretation. S.A.; Was involved in conducting the experiments, manuscript proof, administrative and financial support. All authors read and approved the final manuscript.

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Investigation of Signals and Transcription Factors for The Generation of Female Germ-Like Cells

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Abstract

Objective: Primordial germ cell (PGCs) lines are a source of a highly specialized type of cells, characteristically oocytes, during female germline development in vivo. The oocyte growth begins in the transition from the primary follicle. It is associated with dynamic changes in gene expression, but the gene-regulating signals and transcription factors that control oocyte growth remain unknown. We aim to investigate the differentiation potential of mouse bone marrow mesenchymal stem cells (mMSCs) into female germ-like cells by testing several signals and transcription factors.

Materials and Methods: In this experimental study, mMSCs were extracted from mice femur bone using the flushing technique. The cluster-differentiation (CD) of superficial mesenchymal markers was determined with flow cytometric analysis. We applied a set of transcription factors including retinoic acid (RA), titanium nanotubes (TNTs), and fibrin such as TNT-coated fibrin (F+TNT) formation and (RA+F+TNT) induction, and investigated the changes in gene, MVH/DDX4, expression and functional screening using an in vitro mouse oocyte development condition. Germ cell markers expression, (MVH/DDX4), was analyzed with Immunocytochemistry staining, quantitative transcription-polymerase chain reaction (RT-qPCR) analysis, and Western blots.

Results: The expression of CD was confirmed by flow cytometry. The phase determination of the TNTs and F+TNT were confirmed using x-ray diffraction (XRD) and scanning electron microscope (SEM), respectively. Remarkably, applying these transcription factors quickly induced pluripotent stem cells into oocyte-like cells that were sufficient to generate female germ-like cells, growth, and maturation from mMSCs differentiation. These transcription factors formed oocyte-like cells specification of stem cells, epigenetic reprogramming, or meiosis and indicate that oocyte meiosis initiation and oocyte growth are not separable from the previous epigenetic reprogramming in stem cells in vitro.

Conclusion: Results suggested several transcription factors may apply for arranging oocyte-like cell growth and supplies an alternative source of in vitro maturation (IVM).

Keywords: Cell Differentiation, Germ cells, Transcription Factors


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Introduction

All infertile couples generate at least one meiotically incomplete oocyte, approximately 7 to 16% (1, 2). Oocyte maturation failure, a bad egg syndrome, is occasionally absolute means no mature oocytes are generated. The key clinical features associated with this syndrome include: i. Primary infertility, ii. Repetitive generation of mostly immature oocytes, iii. Inability of in vitro maturation (IVM) to stimulate maturation, and iv. Breakdown of fertilization despite intracytoplasmic sperm injection (ICSI) (3).

In oocyte differentiation, oocyte growth and meiosis are two key processes. Meiosis initiation is believed to be regulated mainly by retinol’s production, storage, and metabolism and its metabolite retinoic acid (RA). RA signaling is performed by its target genes, such as Stra8, which is essential for the initiation and progression of meiosis. Intrinsic factors such as MVH provide diploid germ cells ready for meiosis initiation when extracellular signals are received (4).

Because of their characteristic properties, ability to self-renewal, cloning, and differentiation into many different cell types as pluripotency, stem cells have been recommended in biomedical applications. Recently, much progress has been obtained in understanding stem cell biology and our ability to manipulate their proliferation and differentiation to get functional cells (5). Hübner et al. (6) show that mouse embryonic stem cells can develop into oogonia that enter meiosis; Bahmanpour et al. (7) improved the rate of in vitro oocyte differentiation by using bone morphogenetic protein 4 (BMP4) and RA along with ovarian somatic cell co-culture. Similarly, nanomaterials have recently been synthesized, which increases the efficiency of stem cell differentiation and their biomedicine applications, respectively (8, 9).

Growth and differentiation factors can adhere to nano-supplied surfaces and they can be effectively delivered into the culture medium. Therefore, functional scaffolds made of TNT can be used to grow, biocompatibility, differentiate stem cells, and regenerate damaged tissues (10). However, the small scale of nanomaterials alters their physicochemical properties, making their interactions...
toxic to stem cells. Therefore, the determination of nanomaterial bio-interactions is essential to increase the success of medical treatments and enhance the safety of biomedical devices (11).

Fibrin, made from fibrinogen and thrombin, has been introduced as a suitable biological polymer in which tissue engineering applications and growth factors are delivered in cell cultures (12). Significant advantages of fibrin hydrogels include flexibility, low cytotoxicity, and high effectiveness on nanomaterials with homogeneous distribution (13). This characteristic leads to improved cell growth, viability, and differentiation in response to growth factors (14). According to the mentioned characteristics of fibrin, chemical, structural and mechanical, a 3D scaffold is suitable for later tissue engineering applications (15, 16).

Here, we applied a set of transcriptional factors, RA and RA+F+TNT formation, comprising the underlying gene regulatory expression and validated these findings with functional screening. Furthermore, we endeavored to reconstitute pluripotent stem cell differentiation and thereby we generated oocyte-like cells competent for oocyte meiosis initiation, oocyte growth, and subsequent fertilization in vitro.

Material and Methods

Nanoparticles synthesis

The TiO2 nanoparticles were synthesized by the hydrothermal method described by Khoshnood et al. (17), then particles from 10 to 15 nm, which present micropores and mesopores on their surface, were obtained.

Isolation and culture

In this experimental study, the flushing technique was performed to isolate mouse bone marrow mesenchymal stem cells (mMSCs) from mice femur bone. The cell suspensions were transferred into 15 mL centrifugation tubes and were resuspended in DMEM medium (F12: REF:32500-035 Gibco, USA) supplemented with 10% FBS (Sigma-Aldrich, USA), 1% penicillin, and streptomycin (BI-1203 BIO IDEA Company, USA) mixture, and 1% Gluta MAX (Gibco, USA), and seeded in 25 cm² culture medium flask for maintaining at 37°C humidified incubator with 5% CO₂. At 80-90% confluency, cells were harvested with 0.05% Trypsin-EDTA solution (Gibco, USA) and replated in treatment groups (RA and RA+F+TNT formation). All protocols followed for the utilization of animals were approved by the Ethics Committee of Islamic Azad University, Science and Research Branch, Tehran, Iran, approval ID: IR.IAU. SRB.REC.1400.276.

Induction of stem cells into female germ-like cells

The cells obtained from the 3rd passage were used for signals and transcription factors to generate female germ-like cells. Then they were seeded at a density of 2×10⁴ per well in 24-well plates and treated with 10³ M RA and 50 µg/ml TNT-coated fibrin (fibrinogen+thrombin 1:1) in the medium as mentioned above for 14 days. The Cells were observed for morphological changes during 14 days of induction, after which immunocytochemistry and quantitative transcription-polymerase chain reaction (RT-qPCR) were performed.

Morphology characterization of F+TNT formation

The phase characterization of TiO2 nanoparticles was determined by X-ray diffraction pattern (XRD) (Model PW1730, PHILIPS, Cu LFF lamp λ=1.540598 A, phase size=0.05°, phase time=1 second, voltage 40 kV, current 30 mA, And 40 mV). The 30λ TNT was then added by incubation at 50 µg/ml (18-20) of TNT-PBS-solution (21) on fibrin to measure their excess biological function and differential behavior. Figures analyzed with scanning electron microscope (SEM) and image j software (v. 1.52).

Flow cytometry analysis

For flow cytometric analysis of cluster-differentiation (CD) mesenchymal superficial markers, after four passages, 2×10⁴ cells were removed for a panel of mMSCs antibodies. The Cells were dispersed with 0.25%- trypsin-EDTA (Gibco, USA) and resuspended in PBS supplemented with 0.5% FBS. The cells were aliquoted into several parts and incubated at 4°C for 20 minutes in the dark with monoclonal antibodies (AB92574, AB114052, AB28364, and AB10558) against the hematopoietic cell markers CD31-PE and CD45-PE, and CD90-PE, and CD105-FITC (AB6785 and AB6717), MSC markers CD90-PE, and CD105-FITC (AB6785 and AB6717). Negative control samples were incubated with mouse IgG1- FITC/PE (11-4724) isotype antibodies to help differentiate nonspecific background signals from specific antibody signals. The samples were analyzed on a Partec cytometer (German), and the resulting data were processed using FloMax software (22).

Immunocytochemistry

For Immunocytochemistry analysis of the specific germ cell marker, MVH, the cells were washed with PBS after 14 days of induction and were fixed in 4% paraformaldehyde for 20 minutes at 25°C. The cell membrane was permeabilized with 0.1% Triton X-100 solution in PBS for 20 minutes. Nonspecific binding-site blocking was performed with 5% goat serum for 45 minutes without washing, then incubation with anti-MVH (Mouse monoclonal anti-human, 1:100; Abcam, USA) antibody overnight at 4°C. Subsequently, the cells were washed with PBS and incubated with FITC-conjugated Goat Anti-Rabbit (1:100; Abcam, USA) or Goat Anti-Mouse (1:100; Abcam, USA) for 1 hour at 25°C. Finally, the Nuclei were counterstained with DAPI (Sigma, UK) for 5 minutes, and an Immunofluorescence image was taken using a fluorescent microscope (22).
Quantitative transcription-polymerase chain reaction

Using Tri-Pure reagent (Invitrogen, San Diego, CA, USA) whole RNA isolate per the manufacturer’s instructions. The DNA contamination in the RNA sample was deleted by RNase-free DNase I (Thermo Scientific, USA) for 30 minutes at 37°C. The RNA concentration and purity were specified using the spectrophotometric (WPA spectrophotometer, Biochrom, UK) method. Using a Transcriptor First Strand cDNA Synthesis kit (Roche), the RNA was reversely transcribed by random Hexamer and 1000 ng of DNA-free RNA. TaqMan probe (Life Technologies, India) was applied to survey the expression of MVH, which normalized against 18 seconds expression as a housekeeping gene β-actin. The PCR reaction components were mixed to procure a final volume of 20 μL. The following components were applied: 0.5 μL (25 ng) cDNA, 1 μL TaqMan assay reagent, 10 μL TaqMan universal master mix, and 8.5 μL distilled water. The PCR cycling was as follows: 10 minutes at 95°C, polymerase activation, 40 cycles at 95°C for 15 seconds, and 60°C for 1 minute using a Rotor-Gene Q instrument (Qiagen, Germany). The relative expression of the gene, using the ∆∆Ct method, was analyzed by normalizing the Ct values of the target against 18 seconds (22). Sequences of the MVH primers used for RT-qPCR are:

F: 5’GTGGAAGTGGTCGAGGTGGT3’
R: 5’CTGGTGGAGGAGGGGGTA3’

and primers sequences of housekeeping gene, β-actin are:

F: 5’TCAGAGCAAGAGAGGCATCC3’
R: 5’GGTCATCTTCTCACGGTTGG3’

Western blots

Fifty μg of proteins extracted by RIPA lysis buffer was used for sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to nitrocellulose membranes. The membrane was stained with rabbit polyclonal antibody against MVH (AB13840, Abcam, UK, 1:100 dilutions), and goat anti-mouse secondary antibodies (ab6789, Lucerna-Chem, Switzerland). The antibody-antigen interaction in the membranes was observed using an enhanced-chemiluminescent detection kit (Santa Cruz Biotechnology Inc, Santa Cruz, CA, USA) (23).

Statistical analysis

All data were presented as mean ± SD (standard deviation) and were analyzed using SPSS (v 23, IBM, USA). One-way ANOVA and Tukey tests followed by Bonferroni posttest were used to compare the groups. The significant differences (P<0.05) were calculated among various treatment groups.

Results

Flow cytometry

We evaluated several stem cell-associated CD markers using flow cytometry to determine the CD markers of cultured cells in passage 3. The results illustrate the differential characteristics of stem cells and rule out the hematopoietic origin of isolated cells. The Cells were mostly positive for mMSCs markers, CD90 and CD105, and at least reacted with hematopoietic markers, CD31 and CD45: 95% of the cells express CD90, 72.6% of the cells express CD105, 0.23% of the cells express CD31, and 1.42% of the cells express CD45 (Fig.1). Negative control samples were incubated with IgG1-FITC/PE isotype antibodies in mice. The data represented three independent experiments as mean ± SD.

Structural determination of TNTs and F+TNT formation

The XRD data for the sample synthesized at 120°C is consistent with the standard Anatase (USnano-America) pattern. Usually, diffraction peaks at 30°C indicate the presence of crystal defects or long-term non-sequence in TiO2 nanoparticles. However, such a peak was not observed in our diffraction pattern which suggests the pure anatase phase of TiO2 is formed by a quadrilateral anatase structure at 120°C. The absence of diffraction peaks at 27°C and 31°C indicates that this sample was free of rutile and TiO2 brookite structures. The XRD pattern showed anatase phase TiO2 nanoparticles only by diffraction at angles of 25, 37, 48, 54, 55, 63, 69, and 75°C (peak A, Fig.2A). The above results confirmed that the optimal calcination temperature for preparing pure anatase of TiO2 nanocrystals by polymer gel was 120°C. The morphology of F+TNT formation was confirmed using SEM. It was observed that the surface area of TNT was increased compared with self-aggregation-TNT (Fig.2B).
Immunocytochemistry staining for specific oocyte-like cell marker, MVH, differentiation

The germ cell-related gene, MVH, was detected in all isolated cells using immunocytochemical analysis after 14 days. This experiment confirms that all mouse bone marrow cells have a differentiation potential into germ- and oocyte-like cells with signals and transcription factors. Under immuno-fluorescence microscopy, green fluorescent protein (GFP) expression was highly observed in mMSCs-derived female germ-like cells, treated with RA+F+TNT or RA compared to control cells. (Fig.3A, B).

RT-qPCR analysis of specific oocyte-like cell marker, MVH

RT-qPCR results show that all the cells expressed oocyte-like cell genes after 14 days of transcription factor inductions. However, the quality and quantity of expression differed among two germ cell marker groups, MVH. In this experiment, F+TNT transcription factors showed more significant potential to be differentiated into oocyte-like cells. A significantly higher relative gene expression level of F+TNT was observed compared with RA and control (P<0.05). The results of this analysis show that co-administration of transcription factors has a higher potential for differentiation of mMSCs into female germ-like cells than individual transcription factor (Fig.4).
Western blot analysis measured the expression of female germ cell-associated proteins (MVH) during oocyte-like cells formation from mMSCs after being cultured for 14 days. Furthermore, the level of MVH protein was higher in mMSCs treated with RA or RA+F+TNT compared to the level of MVH protein in the control cells (Fig.5).

**Fig.5:** Western blot analysis of MVH expression in mice bone marrow-derived mMSCs. A. MVH protein levels were evaluated as specific differentiation markers to compare GAPDH reference protein using western blot analysis. B. Protein lysates from mMSCs were blotted and stained by MVH antibody. The level of MVH protein expression was higher in mMSCs treated with RA or RA+F+TNT compared to the level of MVH protein expression in the control cells. **; P<0.0034, ****; P<0.0001, indicated significant differences in the MVH expression between RA+F+TNT, RA, and the control, mMSCs. Mouse bone marrow mesenchymal stem cells, RA; Retinoic acid, F; Fibrin, and TNT; Titanium nanotube.

**Discussion**

This study defines a set of transcription factors, which are promoted the differentiation of oocyte-like cells from mMSCs in vitro, which is expressed germ-like cell marker MVH/DDX4 at both mRNA and protein levels. This study will be a powerful protocol to elevate our understanding of the mechanisms underlying reprogramming in oocyte-like cell growth and of their potential application in IVM technologies.
Oocyte differentiation and development depend on continuous signaling interactions with somatic follicle cells in vivo (24). Signaling molecules; maturation-promoting factor (MPF), transcription, and translation of critical regulatory enzymes are the processes by which an oocyte acquires meiotic competence in vivo. The complex interaction between these factors and the determination of arrest versus progression is related to the delicate balance between the production and targeted degradation of signaling molecules, MPF (25-31). Finally, dysfunction on the molecular level, errors in MPF, and aberrations in chromosomal/spindle formation lead to meiotically oocyte maturation arrest. In the case of reports, changes in gonadotropin stimulation protocol, using IVM, and ICSI does not improve the treatment outcome for bad egg syndrome (3).

RA has been proven to act as a meiosis-inducing factor of meiosis in mouse gonads. RA induces Stra8, an RA-responsive gene in female primordial germ cell (PGCs), leading to meiosis in fetal female germ cells. Recently, reports have shown that RA can induce meiosis in PGCs before gonadal sex differentiation (32-37).

This study introduced several transcription factors, RA, RA+F+TNT, that can induce mMSCs into oocyte-like cells in vitro. Growth and differentiation factors can adhere to nanomaterial and fibrin, which enhance their effective delivery in culture medium and increase the developmental levels. The differentiated oocyte-like cells indicated that transcription factors are a prerequisite to activating the gene-excitatory expression driving oocyte growth in vitro. As such, our results complement recent studies which suggest that the principal role of epigenetic reprogramming is to activate the meiotic program. In addition, our observation showed that oocyte-like cell induction during oocyte growth is dependent on the epigenetic reprogramming in female germ-like cells. This finding provides new insight into the importance of epigenetic control and transcription factors in oocyte maturation. Furthermore, understanding possible roles of transcription factors in oocyte-like cell induction is necessary for new research. The culture conditions are a unique material that is invaluable for applications in assisted reproductive technology, such as IVM and ICSI.

Oocyte-like generation is challenging in vitro, and this involves both cytoplasmic and nuclear processes. This study showed the increased MVH expression at both mRNA and protein levels, as a result of using several transcription factors, RA, TNT, and fibrin, which indicates the effectiveness of these transcription factors in the production of female germ-like cells.

The mMSCs can be reprogrammed to an oocyte-like cell by transcription factors, RA, RA+F+TNT formation; little is known about factors that induce this reprogramming in vivo and in vitro. Furthermore, by understanding the mechanism of oocyte-like cell maturation in vitro, it is possible that IVM protocols could be promoted to obtain the signaling and transcription factors necessary for oocyte maturation competence and performance in vitro.

Conclusion
Here, we demonstrate the induction of pluripotent stem cells from mouse bone mMSCs by introducing germ factors, MVH, under stem cell culture conditions and morphology determination of F+TNT induction. mMSCs exhibit the morphology and growth properties of germ-like cells and express germ cell marker genes following treatment with the above-mentioned transcription factors. These data demonstrate that oocyte-like cells can be directly generated from mMSCs by adding only a few defined factors.

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Authors’ Contributions
S.E.; Conceived the presented idea. S.E., A.Sh., P.M., A.H.; Developed the proposal and performed the experiments. P.M., A.H.; Verified and monitored the analytical methods. S.E., A.Sh.; Investigated morphology characteristics of the TNTs and the TNTs-coated fibrin. A.Sh., P.M., A.H.; Supervised the findings of this work. All authors read and approved the final manuscript.

References
MSCs-Derived Female Germ-Like Cells


Endothelin-1 Stimulates PAI-1 Protein Expression via Dual Transactivation Pathway Dependent ROCK and Phosphorylation of Smad2L

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Abstract

Objective: In addition to the carboxy region, Smad2 transcription factor can be phosphorylated in the linker region as well. Phosphorylation of Smad2 linker region (Smad2L) promotes the expression of plasminogen activator inhibitor type 1 (PAI-1) which leads to cardiovascular disorders such as atherosclerosis. The purpose of this study was to evaluate the role of dual transactivation of EGF and TGF-β receptors in phosphorylation of Smad2L and protein expression of PAI-1 induced by endothelin-1 (ET-1) in bovine aortic endothelial cells (BAECs). In addition, as an intermediary of G protein-coupled receptor (GPCR) signaling, the functions of ROCK and PLC were investigated in dual transactivation pathways.

Materials and Methods: The experimental study is an in vitro study performed on BAECs. Proteins were investigated by western blotting using protein-specific antibodies against phospho-Smad2 linker region residues (Ser245/250/255), phospho-Smad2 carboxy residues (465/467), ERK1/2 (Thr202/Thr204), and PAI-1.

Results: TGF (2 ng/ml), EGF (100 ng/ml) and ET-1 (100 nM) induced the phosphorylation of Smad2L. This response was blocked in the presence of AG1478 (EGFR antagonist), SB431542 (TGFR inhibitor), and Y27632 (Rho-associated protein kinase (ROCK) antagonist). Moreover, ET-1-increased protein expression of PAI-1 was decreased in the presence of bosentan (ET receptor inhibitor), AG1478, SB431542, and Y27632.

Conclusion: The results indicated that ET-1 increases the phosphorylation of Smad2L and protein expression of PAI-1 via induced the transactivation pathways of EGF and TGFR. This study is the first attempt to scrutinize the significant role of ROCK in the protein expression of PAI-1.

Keywords: Atherosclerosis, ROCK, Smad2, Transactivation

Introduction

Endothelin-1 (ET-1) is a strong vasoconstrictor peptide that is synthesized by endothelial cells, probably causing the promotion of endothelial dysfunction (1-4). The effect of ET-1 is exerted through G-protein-coupled receptors (GPCRs): ETₐ and ETₐ (5). GPCR family is the biggest group of cell surface receptors participating in a number of physiological or pathological circumstances (6). Therefore, understanding the different dimensions of GPCR signaling is essential for therapeutic purposes. GPCRs-driven signaling pathways include the classic pathway via direct binding of ligand to GPCRs on the cell membrane leading to activation of heterotrimeric G proteins and multiple signaling pathways. In recent years, transactivation pathways of protein tyrosine kinase receptors (PTK) such as epidermal growth factor receptor (EGFR), as well as protein serine/threonine kinase receptors (PS/TK) like transforming growth factor receptor (TGFR) have been identified as part of the GPCR signaling (7-9). Recent studies have demonstrated that different GPCR agonists such as thrombin, ET-1, and AngII can contribute to transactivation of EGFR and TGFR (10-12). According to our previous study, it has been determined that ET-1 results in TGFR transactivation endothelial cells (13).

TGFB receptors are a group of serine/threonine kinase receptors whose biological roles are performed by type I and type II receptor complexes (ALK5). TGFB signaling is launched by interaction of a ligand to the TβRII/type I heterogenic complex leading to phosphorylation of the carboxy region of Smad proteins (14). Smad proteins are transcriptional factors that play a serious role in the TGFB-superfamily signals (15, 16). The Smads have three distinct regions: two conserved regions including N-terminal (MH1) and C-terminal (MH2) regions, and one non-conserved region -linker region- that links MH1 and MH2 regions. Besides the carboxy region, the linker region can be phosphorylated as well (16-18). In the Smad-dependent TGFB signaling pathway, phosphorylation of C-terminal region occurs immediately by binding of TGFB to the cell surface receptor. However,
in non-Smad signaling, phosphorylation of Smad2 linker region (Smad2L) occurs indirectly by an activated serine/threonine kinase such as ERK1/2, p38, or JNK. Recent studies have shown that in addition to TGF-β, GPCR agonists result in phosphorylation of Smad2L which can play a significant part in regulation of Smad’s function (14). Phosphorylation of Smad2L increases the expression of proteoglycan synthesizing genes. It has been demonstrated that TGF-β/Smad pathway increases plasminogen activator inhibitor type 1 (PAI-1) expression in different cell types (19, 20). PAI-1 is a member of the superfamily of serine-protease inhibitors (serpin) that may cause vascular disorders such as endothelial dysfunction (21, 22). Studies have shown that growth factors such as TNF-α, TGF, GPCR agonists such as thrombin, and angiotensin II can lead to increased mRNA expression of PAI-1 (16, 21, 23). In 1996, it was shown for the first time that angiotensin II (Ang II) can induce transactivation pathways. Subsequently, some comprehensive researches have focused on understanding the underlying mechanism of transactivation pathway in different cell types. However, the details of this pathway and the signaling molecules that participate in transactivation pathways induced by ET-1 are not very clear in bovine aortic endothelial cells (BAECs). Therefore, in the current study and for the first time, not only the role of dual transactivation pathways induced by ET-1 were evaluated in phosphorylation of Smad2L and PAI-1 expressions in BAECs, but also the role of ROCK assessed in the ET-1 induced PAI-1 expression.

Materials and Methods

This experimental study was approved by the Ethics Committee of Ahvaz Jundishapur University of Medical Sciences (IRAJUMS.REC.1396.1.4). Fetal bovine serum (FBS), penicilllin-streptomycin solution, and low glucose (1 g/l) Dulbecco’s modified Eagle’s medium (DMEM) were obtained from Gibco (Invitrogen, Carlsbad, CA, USA). EGF, ET-1, Y27632, AG1478, SB431542, and neomycin were purchased from Sigma-Aldrich (St. Louis, MO, USA). Recombinant transforming growth factor-β, HRP anti-rabbit IgG-peroxidase antibody produced in goat, anti-phospho-Smad2L (ser245/250/255) rabbit polyclonal antibody, anti-phospho-Smad2C (ser465/467) rabbit polyclonal antibody, anti-phospho-ERK1/2(The202/204), PAI-1 antibody, and GAPDH were purchased from Cell Signaling Technology (Beverly, MA, USA).

Cell culture

Bovine aortic endothelial cells (BAEC) were gifted by Professor Peter J Little (School of Pharmacy, The University of Queensland, Australia). BAECs were cultured according to a previously-described procedure (13). In brief, the cells were cultivated in DMEM with 1 g/l glucose containing 10% FBS and 1% antibiotic; and when cells reached about 80 % confluence, they were pretreated with specific inhibitors in certain intervals. In the next step, ET-1 was added to the culture medium. BAECs were incubated with TGFβ (2 ng/ml) for 1 hour and with EGF (100 ng/ml) for 5 minutes, once alone and once in combination with each other (13, 24). To investigate the effects of ET-1 on phosphorylation of Smad2L, the BAECs were treated with ET-1 (100 nM), and then harvested at 5 and 15 minutes, 2, 4, and 8 hours intervals. In order to evaluate Smad2C phosphorylation, BAECs were treated with ET-1 (100 nM) and harvested at 1, 2, and 4 hours intervals (13). In order to evaluate ERK phosphorylation, BAECs were treated with ET-1 (100 nM) and were subsequently harvested at 5, 15 and 30 minutes, 1, 2, 4, and 8 hours intervals (25). The effects of SB431542 (10 μM for 30 minutes) and AG1474 (10 μM for 30 minutes) (24) inhibitors on pSmad2L were tested by pretreating the cells with them. Thereafter, ET-1 (100 nM) was added to the culture medium. The neomycin (100 μM for 1 hour) (26) and Y27632 (10 μM for 30 minutes) (13) inhibitors were tested on pSmad2L via pre-incubation of the cells prior to addition of ET-1 (100 nM) to the culture medium. To investigate the effects of ET-1 on protein expression of PAI-1, the BAECs were treated with ET-1 (100 nM) and then harvested at 30 minutes, 1, 2, 4, and 8 hours intervals (13). The effects of SB431542 (10 μM for 30 minutes) (18) and AG1474 (10 μM for 30 minutes) (24) inhibitors on protein expression of PAI-1 were tested by pretreating the cells with them prior to adding ET-1 (100 nM) to the culture medium. The neomycin (100 μM for 1 hours) (26) and Y27632 (10 μM for 30 minutes) (13) inhibitors were tested on protein expression of PAI-1 via preincubation of the cells prior to addition of ET-1 (100 nM) to the culture medium. The cells were harvested after 4 hours.

Western blot

Proteins were determined using the method of Seif et al. (4). Briefly, harvested cells were lysed in RIPA buffer. Then, the proteins were separated on 10% SDS-PAGE and transferred to a membrane (PVDF). After blocking steps, the membranes were incubated with primary antibodies. The membranes were washed and then exposed with a secondary anti-rabbit IgG antibody conjugated to horseradish peroxidase. The labeled antibodies were detected with chemiluminescence exposure.

Statistical analysis

The results are presented as mean ± SEM of three individual experiments. Statistical significance was estimated by one-way ANOVA, followed by the least significant difference post-hoc analysis (LSD). P<0.05 or P<0.01 considered as statistically significant. Fold change was calculated by dividing all the measured values from the intensity of each area by their controls (for both target and internal control). The areas were obtained using Image J software program. Then, the values of target groups were divided by the values of their control. Graph Pad Prism software program was used for drawing the graphs.
Results

TGF and EGF induced Smad2L phosphorylation in BAEC

To investigate the role of TGFβ and EGF in phosphorylation of Smad2L (ser245/250/255), BAECs were incubated with TGFβ (2 ng/ml) and EGF (100 ng/ml) for 1 hour and 5 minutes, respectively, once alone and once in combination with each other. TGFβ (P<0.05) and EGF (P<0.05) stimulated Smad2L phosphorylation, and the effects of combination of TGF and EGF could be additive to Smad2L phosphorylation (P<0.01, Fig.1). This data demonstrates that both EGF and TGFβ are individually involved in the phosphorylation of Smad2L through two distinct pathways.

ET-1 stimulated Smad2L phosphorylation in BAECs

The effects of ET-1 on phosphorylation of Smad2L (ser245/250/255) were investigated in different times. BAECs were exposed to ET-1 (100 nM) and phosphorylation of Smad2L measured in a period of 5 minutes to 8 hours. ET-1 led to time-dependent increase in Smad2L phosphorylation at hours two (P<0.05) and four (P<0.01, Fig.3A). These results demonstrated that by induction of TGFR transactivation, ET-1 can stimulate Smad2L phosphorylation in BAEC in a time-dependent manner. Moreover, to examine the role of ET-1 in EGFR transactivation, the phosphorylation of ERK1/2 was investigated as the instant downstream mediator of EGFR activation. BAECs were incubated with ET-1 (100 nM) in certain intervals from 5 minutes to 8 hours. ET-1 stimulated ERK1/2 phosphorylation in different points in time between 5 minutes and 1 hours (P<0.01) and between 2 and 8 hours (P<0.05, Fig.3B). These results suggest that by transactivation of EGFR, ET-1 can lead to phosphorylation of ERK1/2 in BAEC.

ET-1 mediated dual transactivation of EGFR and TGFR in BAECs

The question here is that whether there is evidence that the transactivation signaling pathways induced by GPCR are directly involved in phosphorylation of Smad2L? In order to evaluate the role of ET-1 in TGFR transactivation, the phosphorylation of Smad2C (Ser465/467) was investigated as the instant downstream mediator of TGFR activation. BAECs were treated with ET-1 (100 nM) in a period of 1-4 hours. ET-1 led to time-dependent increase in Smad2C phosphorylation at hours two (P<0.05) and four (P<0.01, Fig.3A). These results demonstrated that by induction of TGFR transactivation, ET-1 can stimulate Smad2C phosphorylation in BAEC in a time-dependent manner. Moreover, to examine the role of ET-1 in EGFR transactivation, the phosphorylation of ERK1/2 was investigated as the instant downstream mediator of EGFR. BAECs were incubated with ET-1 (100 nM) in a period from 5 minutes to 8 hours. ET-1 stimulated ERK1/2 phosphorylation in different points in time between 5 minutes and 1 hours (P<0.01) and between 2 and 8 hours (P<0.05, Fig.3B). These results suggest that by transactivation of EGFR, ET-1 can lead to phosphorylation of ERK1/2 in BAEC.
ET-1 Stimulates of PAI-1 via Transactivation Pathways

for 30 minutes prior to treatment with ET-1 (100 nM) for 4 hours. Phosphorylation of Smad2L (ser245/250/255) was markedly alleviated in the presence of AG1478 and SB431542 (P<0.05, Fig.4A). The results of this experiment indicate that ET-1 can induce the phosphorylation of Smad2L via transactivation of EGFR and TGFR. Furthermore, the roles of ROCK and PLC were examined in ET-1-induced phosphorylation of Smad2L (ser245/250/255). Neomycin (100 μM), the specific inhibitor of PLCβ, was used as the downstream mediator of Gαq for 1 hours prior to treatment with ET-1 (100 nM) for 4 hours, and Y27632 (10 μM), a potent inhibitor of ROCK was used as the downstream mediator of G12/13 for 30 minutes prior to treatment with ET-1 (100 nM) for 4 hours. The results of this work showed that Y27632 can reduce Smad2L phosphorylation (P<0.05), but neomycin cannot do the same (Fig.4B). This shows that stimulation of Smad2L phosphorylation by ET-1 is dependent on ROCK activity.

Fig.3: ET-1 leads to phosphorylation of Smad2C and ERK1/2. BAECs were incubated with ET-1 (100 nM). ET-1; Endothelin-1, *; P<0.05, **; P<0.01 vs. untreated, min; Minutes, and h; Hours. Values are presented as mean ± SEM of three individual experiments.

Fig.4: ET-1 leads to phosphorylation of Smad2L via both dual transactivation and the ROCK activity. A. BAECs were preincubated with SB431542 (10 μM) for 30 minutes and AG1478 (10 μM) for 5 minutes before stimulation with ET-1 (100 nM) for 4 hours. B. BAECs were preincubated with neomycin for 1 hours (100 μM) and with Y27632 (10 μM) for 30 minutes before stimulation with ET-1 (100 nM) for 4 hours. ET-1; Endothelin-1, *; P<0.05 vs. untreated, #; P<0.05 vs. ET-1 treated. Values are presented as mean ± SEM of three individual experiments.
ET-1 stimulated the protein expression of PAI-1 in BAECs

The effects of ET-1 were examined on the protein expression of PAI-1. BAECs were exposed to ET-1 (100 nM) at certain points in time from 30 minutes to 8 hours. ET-1 induced the protein expression of PAI-1 at hours one and two (P<0.05) and four (P<0.01, Fig.5). Overall, the results showed that ET-1 increases the protein expression of PAI-1 in BAEC. We chose four hours’ incubation time with ET-1 for the next experiments.

ET-1 stimulates the protein expression of PAI-1 in BAECs through dual transactivation of TGFR and EGFR, as well as the ROCK activity

To assess whether ET-1 leads to an increase in protein expression of PAI-1 via its receptor with induction of dual transactivation, AG1478 (10 μM) and SB431542 (10 μM) as EGFR and TGFR inhibitors, respectively, and Bosentan (10 μM) as ET receptor inhibitor were utilized for 30 minutes prior to treatment with ET-1 (100 nM) for 4 hours. The results showed that ET-1-increased protein expression of PAI-1 was reduced in the presence of AG1478 (P<0.05), SB431542 (P<0.05), and Bosentan (P<0.05, Fig.6A). The present work’s data showed that via its receptor, ET-1 can transactivate EGFR and TGFR in order to stimulate the protein expression of PAI-1. Moreover, in order to study the roles of ROCK and PLC as mediators of the transactivation pathway in protein expression of PAI-1, neomycin (100 μM) for 1 hour and Y27632 (10 μM) for 30 minutes as inhibitors of PLCβ and ROCK were used before being stimulated with ET-1 (100 nM) for 4 hours. The results showed the significant reduction of the protein expression of PAI-1 in the presence of Y27632 (P<0.05), while neomycin could not inhibit the protein expression of PAI-1 (Fig.6B). From this data, it was concluded that ET-1 increased the protein expression of PAI-1, which was dependent on ROCK activity.

Fig.5: ET-1 leads to an increase in the protein level of PAI-1. BAECs were incubated with ET-1 (100 nM) for 30 minutes to 8 hours. Values are presented as mean ± SEM of three individual experiments. ET-1; Endothelin-1, *; P<0.05, **; P<0.01 vs. untreated, min; Minutes, and h; Hours.

Fig.6: ET-1 leads to an increase in the protein level of PAI-1 via induction of dual transactivation pathways, as well as the ROCK activity. A. BAECs were preincubated with SB431542 (10 μM), AG1478 (10 μM), and Bosentan (10 μM) 30 minutes before being stimulated with ET-1 (100 nM) for 4 hours. B. BAECs were preincubated with neomycin (100 μM) for 1 hour and Y27632 (10 μM) for 30 minutes before stimulation with ET-1 (100 nM) for 4 hours. ET-1; Endothelin-1, *; P<0.05 vs. untreated, #; P<0.05 vs. ET-1 treated. Values are presented as mean ± SEM of three individual experiments.
Discussion

In this study, the role of ET-1-induced dual transactivation pathways of EGFR and TGFR were investigated in phosphorylation of Smad2L, as well as the protein expression of PAI-1. Here, it was demonstrated that ET-1 stimulates Smad2L phosphorylation and increases the level of PAI-1 protein through transactivation of EGFR and TGFR, and that ROCK has a central role in this pathway. TGF-β1, alone and in combination with EGF, induced the phosphorylation of Smad2L, which is consistent with the data put out by Kamato et al. (10). Phosphorylation of Smad2L was increased by EGF and TGF-β, indicating the presence of the active pathways of these growth factors in induction of Smad2L phosphorylation. Recent studies have shown that in addition to EGF and TGF-β, GPCR agonists result in activation of kinases such as NOX, P38, and ERK through induction of TGFR and EGFR transactivation (27). Our results showed that ET-1 increased the phosphorylation of Smad2L in BAECs. Kamato et al. (10) presented the evidence that thrombin leads to phosphorylation of Smad2L in VSMCs via transactivation-dependent signaling pathways. In this study, the focus was on the signaling pathways causing the phosphorylation of Smad2L. It was demonstrated that ET-1 stimulated the phosphorylation of Smad2C via TGFR transactivation, as well as ERK1/2 via EGFR transactivation, in BAEC. In a study recently published by the authors, it has been shown that ET-1 stimulates the phosphorylation of Smad2C via TGFR transactivation in BAEC (13). Burch et al. (28) showed that via PAR-1, thrombin can not only lead to EGFR transactivation, but also TGFR transactivation in VSMCs.

It was found that AG1478 (EGFR antagonist) and SB431542 (TGFR antagonist) reduced the effect of ET-1 on phosphorylation of Smad2L, suggesting that ET-1 mediated the phosphorylation of Smad2L through dual transactivation of EGFR and TGFR. Kamato et al. (24) demonstrated that thrombin stimulated the phosphorylation of Smad2L through transactivation of both EGFR and TGFR in VSMC. GPCRs are the biggest cell-surface receptors without any enzymatic activity. These receptors associate with G proteins including Gα, Gβ, and Gγ. Activated G proteins interact with diverse mediators and can regulate signaling responses (8). Signaling pathways that are activated by G proteins are comprised of the following: phospholipase Cβ, adenylate cyclase (AC), and cyclic adenosine monophosphate (cAMP) pathways, as well as Rho kinase (ROCK) (29).

There have been several studies on the roles of these mediators in transactivation pathways. EGFR transactivation is stimulated by Ang II via increasing intracellular Ca²⁺ and activation of PLC/IP3 pathway (30). On the other hand, another study concluded that EGFR transactivation was induced by Ang II, independent of intracellular calcium concentration and PLC/IP3 pathway (31). ROCK signaling leads to transactivation of RSTK in the epithelial cells of mouse lung (32). Therefore, in this study he roles of ROCK and PLC were assessed in Smad2L phosphorylation. It was found that Y27632 (ROCK antagonist) decreased the phosphorylation of Smad2L that was induced by ET-1, but neomycin (PLC antagonist) had no effect on Smad2L phosphorylation. According to the previous studies, ROCK leads to phosphorylation of Smad2C through TGFR transactivation (13, 28, 33). Therefore, based on the results of this study, it is suggested that ROCK has an important role in ET-1 transactivation pathways and subsequently Smad2L phosphorylation. This result is consistent with earlier studies, showing that thrombin stimulated the phosphorylation of Smad2L which is dependent on MMP and ROCK activities in VSMCs (23). Wang et al. (34) showed that PAI-1 is a remarkable prognosticator of cardiovascular disease -dependent death. PAI-1 is a significant factor in the pathophysiology of vascular sclerosis. PAI-1 is mostly expressed by endothelial cells as well as tissues with elevated TGF-β (35). Multiple studies have confirmed that TGF-β1-induced PAI-1 expression occurs via stimulation of EGFR transactivation in vascular, epithelial, and endothelial cells (35, 36). The current study demonstrated that ET-1 increased the level of PAI-1 expression in BAECs in four hours after treatment with ET-1; however, this response was decreased eight hours after the treatment. Therefore, it is possible that deactivation of ET-1 occurred in eight hours. The same pattern can be seen in Smad2L phosphorylation, thus verifying that these pathways are related together. Cell lines alter in morphology, response to stimuli, growth rates, gene and protein expression in different passage numbers (37). The changes observed in the protein expression particularly in the control group in various experiments, may be influenced by different passage numbers, which can be considered as a limitation of this study. In addition, Cockell et al. (38) showed that thrombin induces antigen, natural activity, and mRNA expression of PAI-1 in baboon aortic smooth muscle cells (BASMC). Kerins et al. (39) indicated that Ang IV can stimulate the endothelial expression of PAI-1 via induction of an endothelial receptor.

The protein expression of PAI-1 that is stimulated by ET-1 is decreased in the presence of SB431542 and AG1478. It is suggested that induction of PAI-1 by ET-1 is intervened by transactivation of EGFR and TGFR. Chaplin et al. (33) reported that thrombin transactivated EGFR and TGFR, which can phosphorylate Smad2L and ERK1/2, increase the gene expression of CHSY1 enzymes in VSMCs. The ET-1-stimulated protein expression of PAI-1 was blocked in the presence of ET receptor antagonist (bosentan), strongly suggesting that this response is mediated via the ET-1 receptor. This study has been the first attempt to scrutinize the significant role of ROCK in protein expression of PAI-1 in BAECs. To examine the importance of ROCK and PLC as mediators of activated G proteins, the level of PAI-1 protein was investigated in the presence of Y27632 (ROCK antagonist) and neomycin (PLC antagonist). In a previous study by the authors, it was shown that ET-1 receptor can transactivate the TGFR and then phosphorylate Smad2C.
It was demonstrated that Rho/ROCK kinase plays an important role in mediating the transactivation of TGFR and phosphorylation of Smad2C (Ser465/467) induced by ET-1 (13). Moreover, in another study by the authors that has not yet been published, the role of Rho/ROCK kinase is investigated in ET-1-induced EGFR transactivation. In the current study, a decrease in PAI-1 protein expression was observed in the presence of Y27632 (ROCK antagonist) that could inhibit Smad2L phosphorylation. Based on the results obtained from this study, Rho/ROCK kinase (as a mediator of the transactivation pathway) has a role in Smad2L phosphorylation and PAI-1 protein expression.

In this study, it was shown that ROCK is involved in PAI-1 protein expression for the first time. Observations of the current work strongly suggest that ROCK induced Smad2L phosphorylation via transactivation and affected the enhancement of PAI-1 protein expression. TGF-β1 Smad2L phosphorylation via transactivation and affected the current work strongly suggest that ROCK induced PAI-1 protein expression for the first time. Observations obtained from this study, Rho/ROCK kinase (as a mediator of the transactivation pathway) has a role in Smad2L phosphorylation and PAI-1 protein expression.

The current study demonstrated that ET-1 stimulated the phosphorylation of Smad2L, and this reaction was blocked by AG1478 and SB431542, suggesting that ET-1 leads to Smad2L phosphorylation via induction of dual transactivation of EGFR and TGFR. According to the results of previous studies, ROCK has a key role in inducing transactivation pathways. Hence, induction of Smad2L phosphorylation through dual transactivation of EGFR and TGFR is dependent on ROCK signaling. Furthermore, it was demonstrated that ET-1 increased the level of PAI-1 protein via transactivation of EGFR and TGFR, which is associated with promoting the intravascular thrombosis and atherosclerosis. Moreover, this cellular response is also dependent on ROCK signaling. Therefore, it can be concluded that Smad2L phosphorylation and promotion of PAI-1 protein level may be related together. However, further studies are needed to identify this signalling.

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**Authors’ Contributions**

F.S.; Performed all in vitro experiments, analyzed the data, and wrote the manuscript. H.B.-R., A.Kh.; Contributed to concept and design, financial support, and final approval of the manuscript. All the authors read and approved the final manuscript.

**References**


**MiRNA-16-1 Suppresses Mcl-1 and Bcl-2 and Sensitizes Chronic Lymphocytic Leukemia Cells to BH3 Mimetic ABT-199**

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**Abstract**

**Objective:** Chronic lymphoid leukemia (CLL) is the most common type of leukemia among adults. Increased levels of Mcl-1 and Bcl-xL is linked to resistance to Bcl-2 inhibitors including ABT-199. In this study, we investigated the effect of miRNA-16-1 on apoptosis and sensitivity of the CLL cells to ABT-199.

**Materials and Methods:** In this experimental study, the Mcl-1 and Bcl-2 expression were measured using qualitative reverse transcription-polymerase chain reaction (qRT-PCR) and western blotting. The effect of treatments on cell survival and growth were explored with MTT assay and Trypan blue assay, respectively. The drug interaction was evaluated using combination index analysis. Apoptosis was assessed by ELISA cell death and caspase-3 activity assays.

**Results:** MiRNA-16-1 markedly inhibited the expression of Mcl-1 and Bcl-2 in a time dependent manner (P<0.05, relative to blank control). Pretreatment with miRNA-16-1 synergistically suppressed the cell growth and survival and reduced the half-maximal inhibitory concentration (IC50) value of ABT-199. Moreover, miRNA-16-1 markedly augmented the apoptotic effect of ABT-199 in CLL cells (P<0.05).

**Conclusion:** Our findings propose that miRNA-16-1 act in concert with ABT-199 to exert synergistic anticancer efficacy against CLL, which is attributed to the inhibition of Bcl-2 and Mcl-1. This may propose a promising strategy for CLL resistant patients.

**Keywords:** ABT-199, Bcl-2, Chronic Lymphocytic Leukemia, Mcl-1

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**Introduction**

Chronic lymphoid leukemia (CLL) is the most common type of leukemia among adults with a median age of diagnosis 72 years (1, 2). Despite recent advances in treatments over single-agent chemotherapy such as fludarabine or chemo-immunotherapy combinations, CLL remains an incurable disease (2). A variety of parameters such as deregulated production of survival signals or intrinsic defects in apoptotic machinery contribute to the therapy resistance in CLL patients (3-5). As a result, there remains a need for understanding the detailed molecular pathophysiology of CLL as well as the development of new drugs for clinical treatment of CLL (6).

Apoptosis is induced by the two extrinsic and intrinsic pathways. The intrinsic pathway cell death is controlled by the Bcl-2 family proteins members, including the proapoptotic and the antiapoptotic proteins (5, 7). Overexpression of some antiapoptotic proteins such as Mcl-1 and Bcl-2 is correlated with shorter overall survival and chemoresistance in CLL patients. Accordingly, many targeted strategies have been developed to the Bcl-2 and Mcl-1 to overcome drug-resistance of CLL patients (8, 9).

The Bcl-2-specific antagonist ABT-199 or venetoclax has showed improved clinical efficacy in patients with CLL (10, 11). ABT-199 is demonstrated high cytotoxicity against CLL cells in vitro but is much less effective against CLL cells that have expressed high levels of Mcl-1. Therefore, combination therapy of CLL cell with Mcl-1 inhibitors and ABT-199 have been suggested for improvement of apoptosis-based therapies in CLL resistance cells (12-14).

MicroRNAs (miRNAs) are a small family of endogenous, single-stranded, non-coding RNAs with 20-22 nucleotides in length that are involved in numerous cellular processes such as cell survival, cell death, differentiation and proliferation. They act by directly binding to the specific target mRNA, causing inhibition of the gene expression (15, 16). Several studies have demonstrated that miRNAs are recognized as important diagnostic and therapeutic biomarkers in numerous types of cancers, such as colon and breast cancer (17, 18). Furthermore, miRNAs are involved in almost all hematological processes, suggesting the important role of miRNAs in CLL (19-21). Genetic abnormalities have been observed in the majority of CLL cases. These aberrations include the 11q deletion with intermediate risk, the 13q deletion with low risk, and the 17p deletion with high risk (22). The 13q14 deletion is the most common genetic aberrations observed in more than...
50% of CLL cases. The results of previous studies have clarified that the miRNA-16-1 gene was absent or down-regulated in CLL cases with 13q14 deletion. MiRNA-16-1 is a tumor suppressor gene that involved in the regulation of cell proliferation and cell death via targeting of several molecules (cyclin-dependent kinase 6, cyclin D1, cyclin D3, and Bel-2) (23). In addition, the results of experimental studies show that there is a significant relationship between miRNA-16-1 and Mcl-1 expression levels in samples of CLL patient (19, 24). However, the exact role of miRNA-16-1 in pathogenesis and drug resistance of CLL has not been fully investigated.

We hypothesized that reducing the expression of the miRNA-16-1 gene could lead to increased expression of Bel-2 and Mcl-1, and subsequently resistance to the ABT-199 in CLL cells. Therefore, we investigated the combination effect of miRNA-16-1 and ABT-199 on survival and apoptosis of the CLL cells.

Materials and Methods

Cell culture conditions

The CLL-CII leukemia cells (Pasteur Institute, Iran) were cultured in RPMI-1640 medium (Sigma-Aldrich, St. Louis, MO, USA) containing 10% (vol/vol) heat-inactivated fetal bovine serum (FBS, Sigma Aldrich, USA), streptomycin (100 mg/ml), penicillin (100 U/ml), 1% (v/v) Glutamax (Sigma Aldrich, USA), and 1% sodium pyruvate at 37°C in 5% CO2. The cells were seeded in suspension at a concentration of 1×105 cells/ml with the medium changed every two days.

This research was ethically wise approved from Deputy of Research and Technology, Arak University of Medical Sciences, Arak, Iran (IR.ARAKMU.REC.1395.185).

Transfection of miRNA

The miRNA-16-1 mimics with the sense strand sequence 5'-UAG CAG CAC GUU AAU AUU GGC G-3' and the negative control (NC) miRNA sense strand sequence 5'-ACU ACU GAG UGA CAG A-3' were 5'-ACU ACU GAG UGA CAG UAG A-3' bought from Dharmaco (Lafayette, CO, USA) and used in transient transfection of CLL-II cells. Cell transfection was executed with using Lipofectamine™2000 transfection reagent (Invitrogen, Carlsbad, CA, USA) and Opti-MEM I reduced serum medium (Invitrogen, USA) according to the manufacturer’s recommendation. In brief, the cells were cultured at 40-50% confluence in culture medium without serum and antibiotics one day before transfection. To make the transfection complex, we diluted miRNA-16-1 mimics or NC miRNA (50 nM) with Lipofectamine™2000 (4 µl/ml of transfection medium) in Opti-MEM I. The diluted solutions were thoroughly mixed and incubated for 15-20 minutes at room temperature. Next, the mixture was added to the culture medium. After 6 h of incubation, medium was replaced with a complete growth medium (10% FBS). At different time points after transfection, the CLL-II cells were harvested and various experiments performed.

MTT assay

The cytotoxic effects of miRNA-16-1 and ABT-199 (Sigma- Aldrich, USA) on CLL cells was evaluated using 3-(4, 5-Dimethylthiazol-2-yl)-2, 5-Diphenyltetrazolium Bromide (MTT) assay (25). The experiment was divided into eight groups: ABT-199, miRNA-16-1 mimics, NC miRNA, miRNA-16-1 mimics and ABT-199, NC miRNA and ABT-199, miRNA blank control, ABT-199 blank control and combination blank control. Briefly, the cells were cultivated in 96-well tissue plates at a density of 5×104 cells per well, and then transfected with miRNAs. Six hours later, the cells were treated with different concentrations of ABT-199 (0, 0.05, 0.1, 0.2, 0.4, 0.8, 1.6 and 3.2 µM). After 24 and 48 hours of incubation, the cytotoxicity was determined using a MTT assay kit (Roche Diagnostics GmbH, Mannheim, Germany) according to the manufacturer’s instructions. The absorbance (A) was measured spectrophotometrically at 570 nm with a microplate reader (Awareness Technology, Palm City, FL, USA). Half-maximal inhibitory concentration (IC50) (drug concentration that reduced 50% survival rate) value of the ABT-199, alone or in combination with miRNA, was calculated with Prism 6.01 software (GraphPad Software Inc., San Diego, CA, USA). In the next experiments, the IC50 doses of fludarabine were used.

Combination effect analysis

The combination index (CI) analysis was performed to explore the interaction between ABT-199 and miRNA-16-1 (25-27). The results obtained from the MTT experiment were converted to Fraction affected (Fa, where Fa=0 is 100% cell survival and Fa=1 is 0% cell survival) and analyzed by CompuSyn 1.0 software from Combosyn (Paramus, NJ, USA). Additive, synergistic and antagonistic effects are indicated by CI=1, CI<1 and CI>1, respectively.

Quantitative real time polymerase chain reaction

After treatments, total RNA was extracted by using AccuZol™ reagent (Bioneer, Daejeokgu, Daejeon, Korea) according to the manufacturer’s instructions. Then, reverse transcription of 1 µg of purified total RNA was performed by use of PrimeScript RT reagent kit (Promega, Madison, WI, USA), following the manufacturer’s protocol. Relative gene expression was measured by qualitative reverse transcription-polymerase chain reaction (qRT-PCR) using SYBR Premix Ex Taq (Takara Bio, Otsu, Shiga, Japan) and the LightCycler 96 System (Roche Diagnostics GmbH). RT-PCR carried out in a final volume of 20 µl containing 1 µl of cDNA template, 10 µl of SYBR green reagent and 0.2 µM of each of the primers. The sequences of PCR primers were as follows:
β-actin-
F: 5’-TCC CTG GAG AAG AGC TAC G-3’
R: 5’-GTA GTT TCG TGG ATG CCA CA-3’

MiCl-1-
F: 5’-TAA GGA CAA AAC GGG ACT G-3’
R: 5’-ACC AGC TCC TAC TCC AGC AA-3’

Bcl-2-
F: 5’-ATC GCC CTG TGG ACT GAG T-3’
R: 5’-GCC AGG AGA AAT CAA ACA GAG GC-3’.

The protocol parameters were as follows: initial incubation at 95°C for 3 minutes followed by 40 cycles of denaturation at 95°C for 10 seconds and annealing and extension at 60°C for 50 seconds. The relative mRNA levels were determined using the comparative CT method, 2-ΔΔCt (26, 27), and β-actin as an endogenous control.

Immunoblotting analysis
After treatments, the cells were washed with phosphate-buffered saline (PBS, Sigma-Aldrich, USA) and cell lysates prepared by disrupting cells in lysis buffer (1% NP-40, 0.1% sodium dodecyl-sulfate (SDS, Sigma-Aldrich, USA), 0.5% sodium deoxycholate, 1 mM ethylenediaminetetraacetic acid (EDTA, Sigma-Aldrich, USA), 50 mM Tris·HCl pH=7.4, 150 mM NaCl) containing protease inhibitor cocktail (Roche Diagnostics GmbH). Protein samples (fifty micrograms) were separated on 10% sodium dodecyl-sulfate polyacrylamide gel electrophoresis (SDS-PAGE, Sigma-Aldrich, USA) gels and transferred onto PVDF membrane (GE Healthcare, Amersham, Buckinghamshire, UK). Mouse primary monoclonal antibodies for Mcl-1 (Abcam, Cambridge, MA, UK), Bcl-2 (Abcam) and β-actin (Abcam) were used at 1:1000 dilutions. HRP-conjugated secondary antibodies prepared by disrupting cells in lysis buffer (1% NP-40, 0.1% sodium dodecyl-sulfate (SDS, Sigma-Aldrich, USA), 50 mM Tris·HCl pH=7.4, 150 mM NaCl) containing protease inhibitor cocktail (Roche Diagnostics GmbH). Protein samples (fifty micrograms) were separated on 10% sodium dodecyl-sulfate polyacrylamide gel electrophoresis (SDS-PAGE, Sigma-Aldrich, USA) gels and transferred onto PVDF membrane (GE Healthcare, Amersham, Buckinghamshire, UK). Mouse primary monoclonal antibodies for Mcl-1 (Abcam, Cambridge, MA, UK), Bcl-2 (Abcam) and β-actin (Abcam) were used at 1:1000 dilutions. HRP-conjugated secondary antibodies (Abcam) were used at 1:4000 dilutions. The blot signals were detected using ECL plus western blotting detection Kit (GE Healthcare) and X-ray film (Estman Kodak, Rochester, NY, USA) and quantified via ImageJ 1.62 software (National Institutes’ of Health, Bethesda, Maryland, USA).

Cell growth assay
The effect of miRNA-16-1 and ABT-199 on tumor cell growth was assessed by the trypan blue staining. CLL-CII cells (1×10^5 cells/well) were treated with miRNA-16-1 and ABT-199 in 6-well culture plates for 5 days as described previously. At the end of each day, the cells were collected and cell suspensions stained with 0.4% trypan blue dye (Merek KGaA, Darmstadt, Germany). After 2 minutes of incubation, the number of viable cells was measured using a hemocytometer and an inverted microscope (Nikon Instrument Inc., Melville, NY, USA). The percentage of cell viability in control group was considered as 100%.

Apoptosis ELISA assay
Cell death was determined with an ELISA apoptosis kit (Roche Diagnostics GmbH) that determines monoand oligonucleosomes released into the cytoplasm of apoptotic cells (25). The CLL-CII cells were cultivated at a density of 1×10^5 cells/well in 6-well culture plates and exposed to miRNA-16-1 and ABT-199, as described previously. After 24-48 hours of incubation, the cells were lysed and ELISA assay was performed according to the manufacturer’s instructions. Briefly, 20 µl of the supernatants and 80 µl of immunoreagent containing DNA-peroxidase and histone-biotin antibodies were added to each well of a streptavidin-coated plate and the plate was incubated for 2 hours at room temperature. After washing with incubation buffer, 100 µl of ABTS solution was added. Finally, the reactions were stopped with ABTS stop solution and absorbance was quantified immediately by an ELISA reader (Awareness Technology, Palm City, FL, USA) at 405 nm. Data were calculated as the fold increase in the absorbance of test groups relative to the control group.

Caspase-3 activity assay
The in vitro induction of caspase-3 activity was determined using a colorimetric caspase assay Kit (Abnova, Taipei, Taiwan) (25). Briefly, the treated cells were resuspended in 50 µl cooled lysis buffer and then centrifuged in 10,000 g for 1 minute. Then, 5 µl of the 4 mM DEVD-pNA substrate and 50 µl of 2X reaction buffer (containing 10 mM DTT) were added to each sample. After 2 hours incubation at 37°C the absorbance was quantified using a microplate plate reader (Awareness Technology, Palm City, FL, USA) at 405 nm.

Statistical analysis
All results in this study are demonstrated as mean ± standard deviation (SD) of three experiments. ANOVA followed by Bonferroni’s test was used to determine the significant differences between groups. A P<0.05 was considered significant. All results were analyzed using Prism 6.01 software (GraphPad Software Inc., San Diego, CA, USA).

Results
MiRNA-16-1 inhibited the expression of Mcl-1 and Bel-2 mRNA and protein in CLL-CII cells
First, we explored the effect of miRNA-16-1 on Mcl-1 and Bcl-2 levels in CLL-CII leukemic cells by qRT-PCR and western blotting. As shown in Figure 1A and 1B, transfection of miRNA-16-1 markedly reduced both Mcl-1 and Bcl-2 mRNA levels in a time-dependent way (P<0.05, relative to the blank control). At 24 and 48 hours after treatment with ABT-199, the relative Mcl-1 mRNA expression levels were significantly enhanced, while the expression levels of Bcl-2 mRNA did not change. In miRNA-16-1 and ABT-199 combination group, the expression of Bcl-2 mRNA was similar to the cells transfected with only miRNA-16-1. In addition, the expression of Mcl-1 mRNA in the combination group was
higher and lower than the cells treated with only miRNA-16-1 or ABT-199, respectively. However, NC miRNA had a negligible effect on mRNA expression compared to the blank control (P>0.05). The results of western blotting were in agreement with the PCR results (Fig.1C-F).

Table 1: IC₅₀ of ABT-199 in combination with miRNAs, in CLL cells

<table>
<thead>
<tr>
<th>Treatment</th>
<th>24 hours IC₅₀ (µM)</th>
<th>48 hours IC₅₀ (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABT-199</td>
<td>0.37 ± 1.33</td>
<td>0.24 ± 1.30</td>
</tr>
<tr>
<td>NC miRNA and ABT-199</td>
<td>0.33 ± 1.54⁺</td>
<td>0.22 ± 0.80⁺</td>
</tr>
<tr>
<td>miRNA-16-1 and ABT-199</td>
<td>0.22 ± 2.41⁺</td>
<td>0.14 ± 1.53⁺</td>
</tr>
</tbody>
</table>

IC₅₀ of ABT-199 was calculated by GraphPad Prism 6.01 software and sigmoidal dose-response model. Data expressed as the mean ± SD of three independent experiments. CLL; Chronic lymphoid leukemia,⁺; P<0.05 relative to the corresponding ABT-199, and IC₅₀; Half-maximal inhibitory concentration.

**MiRNA-16-1** synergistically enhanced the effect of ABT-199 on CLL-CII cells

To assess whether the combination of miRNA-16-1 and ABT-199 on CLL-CII cells is synergistic, the combination analysis using the Chou-Talalay method was carried out. The results showed that the effects of miRNA-16-1 (50 nM) and ABT-199 (0.05-3.2 µM) were synergistic with the CI values of >1 in all concentrations of ABT-199 (Fig.2B, D). CI-Fa curved demonstrated that the most synergistic effects of 24 hours (CI=0.77) and 48 hours (CI=0.72) of treatment were seen at 0.4 and 0.2 µM of ABT-199 with Fa values of 0.61 and 0.49, respectively (Table 2).
**Table 2: CI analysis of miRNA-16-1 and ABT-199 in CLL cells**

<table>
<thead>
<tr>
<th>ABT-199 concentration (µM)</th>
<th>24 hours Fa</th>
<th>CI</th>
<th>Combined effect</th>
<th>48 hours Fa</th>
<th>CI</th>
<th>Combined effect</th>
</tr>
</thead>
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<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.05</td>
<td>0.17</td>
<td>0.96</td>
<td>S</td>
<td>0.29</td>
<td>0.78</td>
<td>S</td>
</tr>
<tr>
<td>0.1</td>
<td>0.32</td>
<td>0.89</td>
<td>S</td>
<td>0.41</td>
<td>0.75</td>
<td>S</td>
</tr>
<tr>
<td>0.2</td>
<td>0.47</td>
<td>0.79</td>
<td>S</td>
<td>0.49</td>
<td>0.72</td>
<td>S</td>
</tr>
<tr>
<td>0.4</td>
<td>0.61</td>
<td>0.77</td>
<td>S</td>
<td>0.72</td>
<td>0.73</td>
<td>S</td>
</tr>
<tr>
<td>0.8</td>
<td>0.78</td>
<td>0.81</td>
<td>S</td>
<td>0.81</td>
<td>0.80</td>
<td>S</td>
</tr>
<tr>
<td>1.6</td>
<td>0.86</td>
<td>0.84</td>
<td>S</td>
<td>0.90</td>
<td>0.86</td>
<td>S</td>
</tr>
<tr>
<td>3.2</td>
<td>0.93</td>
<td>0.85</td>
<td>S</td>
<td>0.97</td>
<td>0.91</td>
<td>S</td>
</tr>
</tbody>
</table>

The CI analysis was measured using CompuSyn software and combination index method of Chou-Talalay. CLL; Chronic lymphoid leukemia, CI value >1, =1 and <1 show antagonistic, additive and S effects, respectively, CI; Combination index, S; Synergistic, and Fa; Fraction affected.

**MiRNA-16-1 enhanced the effect of ABT-199 on CLL cell growth**

As over-expression of Mcl-1 and Bcl-2 is linked to the cell growth; we therefore explored whether miRNA-16-1 could inhibit the proliferation of the CLL-CII cells. The CLL-CII cells were treated with miRNA-16-1 (50 nM), ABT-199 (IC50 of 25 hours) and combination of them for 1-5 days and the percent of the viable cells was counted every day by trypan blue staining assay. Data showed that in comparison with the control group, treatment with miRNA-16-1 or ABT-199 significantly suppressed the growth of CLL-CII cells over a period of 5 days. At 24 hours after treatment with miRNA-16-1 or ABT-199, the cell viability dropped to 83.30% and 62.23% respectively, and then to a further 54.15% and 25.67% at the end of the experiment (day 5). Moreover, combination therapy with miRNA-16-1 and ABT-199 had a stronger effect on inhibition of cell growth compared to single therapy (P<0.05). However, no significant difference in cell growth was seen between the NC miRNA and the blank control groups (Fig.3).

**Increased levels of miRNA-16-1 enhanced ABT-199-induced apoptotic**

To explore whether the observed sensitizing effect of the miRNA-16-1 was associated with the increased amount of apoptosis, the effects of miRNA-16-1 and ABT-199 alone and in combination on apoptosis, were assessed using an ELISA apoptosis assay. Results demonstrate that 24 h treatment with miRNA-16-1 or ABT-199 increased apoptosis by 1.93 fold and 4.30 fold, respectively, compared to the control group (Fig.4A, P<0.05). Furthermore, combination treatment enhances the extent of cell death to 7.44 fold (P<0.05), compared with either ABT-199 alone or miRNA-16-1 alone. Moreover, 48 h exposure of the cells with miRNA-16-1 or ABT-199 alone, increased apoptosis by 2.46 and 5.12 fold, respectively, relative to the control group (P<0.05). Also, the combination of miRNA-16-1 and ABT-199 further augmented the induction of apoptosis to 8.56 fold during same period of time (P<0.05, compared with the blank control or monotreatment). However, NC miRNA (alone or in combination with ABT-199) showed no significant effect on extents of apoptosis compared with the miRNA-16-1 or ABT-199, respectively (Fig.4A, P>0.05). The results of ELISA apoptosis assay shows that miRNA-16-1 sensitizes the chronic lymphocytic leukemia cells to ABT-199 partially via enhancement of apoptosis.

**MiRNA-16-1 enhanced the effect of ABT-199 on caspase-3 activity in CLL-CII cells**

To explore the mechanism by which apoptosis occurred...
in the treated cells, changes in the activation of the caspases-3 were determined by using caspase-3 activity assay Kit. Figure 4B shows the changes in caspases-3 activity in the CLL cells treated with the miRNA-16-1, ABT-199 (IC_{50}) and their combination for 24 hours, that show the caspase-3 activity was enhanced by 1.65, 3.42, and 6.21 times, respectively, relative to the blank control group (P<0.05). As indicated in Figure 4B, miRNA-16-1 alone and in combination with ABT-199 activated caspase-3 activity in a time dependent way. However, treatment with NC miRNA did not show a notable effect on caspase-3 activity relative to the blank control group (P>0.05).

**Discussion**

Although ABT-199 has shown high clinical activity against CLL, some patients do not respond or become resistant to this Bcl-2 inhibitor. It has been reported that genetic factors such as Bcl-2 and Bax mutations have been associated with ABT-199 resistance. Sustained activation of B-cell receptor and AKT as well as up-regulation of Mcl-1 and Bcl-xL levels is also related to this process (12-14). However, the exact mechanisms of resistance are not fully known. Our findings propose that miRNA-16-1 act in concert with ABT-199 to exert synergistic anticancer efficacy against CLL, which is attributed to the inhibition of Bcl-2 and Mcl-1.

Our study demonstrated that inhibition of Mcl-1 and Bcl-2 by miRNA-16-1 was associated with inhibition of cell proliferation and increased the sensitivity of the CLL cells to ABT-199 in a synergistic way. So far, various studies have investigated the role of apoptotic proteins, especially Mcl-1, in the sensitivity of tumor cells to ABT-199. For example, Wang et al. (28) showed that Mcl-1-dependent AML cells were resistant to ABT-199 and Mcl-1-specific inhibitors such as A-1210477 that could counteract these resistance in vitro and in vivo. Other study indicated that treatment with A-1592668, a small-molecule inhibitor of CDK9, resulted in the loss of Mcl-1 expression and apoptosis in Mcl-1 dependent lymphoma and AML cell lines. Moreover, the A-1592668 plus ABT-199 combination showed efficacy superior to either agent alone with minimal toxicity in mouse models (29). In addition, Choudhary et al. (13) explored the mechanisms of resistance to ABT-199 in CLL and non-Hodgkin lymphoma cell lines. Their study demonstrated persistent activation of AKT as well as over-expression of Bcl-xL and Mcl-1 levels in the acquired and inherent ABT-199 resistant cells. Moreover, treatment with specific inhibitor of AKT pathway reduced Mcl-1 levels and sensitized the tumor cells to ABT-199. However, our data further confirms the results of the above studies and suggests that downregulation of Mcl-1 by miRNA-16-1 can enhance the ABT-199 sensitivity in CLL cells that depend on Mcl-1 for survival.

MiRNA-16-1 acts as a tumor suppressor by targeting critical molecules in CLL cells. However, few studies have been performed on the role of this miRNA in the chemoresistance of CLL (30). In our study, transfection of miRNA-16-1 increased the ABT-199 sensitivity of the CLL cells. So far, several investigations have been performed to show the relationship of miRNA with chemoresistance. Zhu et al. (31) reported that miRNA-15a, miRNA-16-1, miRNA-34 and miRNA-181a/b sensitized the CLL cells to fludarabine-induced apoptosis through the inhibition of Mcl-1 and Bcl-2. Some other studies of miRNA expression reported that miRNA-221 and miRNA-222a strongly down-regulated in fludarabine-resistant cells in vitro (24, 32). Since the increased expression of Mcl-1 is associated with resistance of tumor cells to Bcl-2-specific inhibitors, other investigations have been performed to
explore the effect of miRNAs on Mcl-1 expression and the sensitivity of tumor cells to these inhibitors. MiRNA-193b is down-regulated in melanoma cells, and induced expression of this miRNA restores ABT-737 sensitivity of the resistant cells by targeting Mcl-1 (33, 34). Similarly Lam et al. (34) recognized a panel of 12 miRNAs that were linked to the reduced Mcl-1 protein levels that can sensitize melanoma cells to the apoptosis induced by ABT-263. In accordance with the above reports, our findings showed that miRNA-16-1 increases ABT-199 sensitivity of the CLL cells. No other study has been done on the relationship between miRNAs and sensitivity to ABT-199 in cancer cells.

We also examined the effects of miRNA-16-1 and ABT-199 on cellular apoptosis. Our results demonstrated that ABT-199 significantly triggered apoptosis and enhanced caspase-3 activity in CLL cells. Moreover, suppression of Mcl-1 and Bcl-2 by miRNA-16-1 was associated with the induction of apoptosis and enhancement of the ABT-199-mediated apoptosis. The intrinsic pathway of apoptosis is induced with different stimuli such as DNA damage, oxidative stress, cytotoxic drugs and radiation. This pathway is under the control of Bcl-2 family of pro- and anti-apoptotic proteins (35, 37). The pro-apoptotic members Bcl-2 family such as Bak and Bax when activated lead to protein hemodimerization, change in the mitochondrial outer membrane permeability (MOMP), release of cytochrome c, and ultimately the downstream activation of the caspases 3, 6 and 7. The anti-apoptotic members such as Bcl-2 and Mcl-1 inhibit apoptosis by heterodimerising with Bak and Bax (5, 9). ABT-199 induces intrinsic pathway of apoptosis in CLL cells by inhibiting Bcl-2. It has been shown that caspase-3 activation is induced by the ABT-199. Moreover, it has been shown that the tumor cells which over-expressed Bcl-2, Mcl-1 and Bcl-xL were resistant to ABT-199 (9, 28, 29). The above reports are in accordance with our results and propose that targeting of anti-apoptotic family members would be a promising strategy to enhance the activity of ABT-199 in various malignancies including CLL.

Conclusion

The data presented here indicate that miRNA-16-1 act in concert with ABT-199 to exert synergistic anticancer efficacy against CLL, attributed to the inhibition of Bcl-2 and Mcl-1. Moreover, our study demonstrated that miRNA-16-1 could augment the execution of apoptosis induced by ABT-199. The intrinsic pathway of apoptosis and caspase activation may be a part of the underlying mechanisms involved in this process. Collectively, our findings show that the combination of miRNA-16-1 and ABT-199 can efficaciously induce the apoptosis of CLL cells, and may offer a promising strategy for patients with CLL.

Acknowledgments

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Authors’ Contributions

H.K.; Study concept and design. N.A., A.S.A.M., M.B.; Acquisition of data. N.A., H.K., M.B., A.S.A.M.; Analysis and interpretation of data. N.A., H.K., M.B.; Drafting of the manuscript. N.A., H.K., A.S.A.M.; Critical revision of the manuscript for important intellectual content. H.K., A.S.A.M.; Funding recipients. All authors read and approved the final manuscript.

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MiRNA-16-1 Sensitizes Leukemia to ABT-199

Genetic and Epigenetic Evaluation of Human Spermatogonial Stem Cells Isolated by MACS in Different Two and Three-Dimensional Culture Systems

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Abstract
Objective: Epigenetic and genetic changes have important roles in stem cell achievements. Accordingly, the aim of this study is the evaluation of the epigenetic and genetic alterations of different culture systems, considering their efficacy in propagating human spermatogonial stem cells isolated by magnetic-activated cell sorting (MACS).

Materials and Methods: In this experimental study, obstructive azoospermia (OA) patient-derived spermatogonial cells were divided into two groups. The MACS enriched and non-enriched spermatogonial stem cells (SSCs) were cultured in the control and treated groups; co-culture of SSCs with Sertoli cells of men with OA, co-culture of SSCs with healthy Sertoli cells of fertile men, the culture of SSCs on PLA nanofiber and culture of testicular cell suspension. Gene-specific methylation by MSP, expression of pluripotency (NANOG, C-MYC and OCT-4), and germ cells specific genes (Integrin α6, Integrin β1, PLZF) evaluated. Cultured SSCs from the optimized group were transplanted into the recipient azoospermic mouse.

Results: The use of MACS for the purification of human stem cells was effective at about 69% with the culture of the testicular suspension, being the best culture system. Upon purification, the germ-specific gene expression was significantly higher in testicular cell suspension and treated groups (P≤0.05). During the culture time, gene-specific methylation patterns of the examined genes did not show any changes. Our data from transplantation indicated the homing of the donor-derived cells and the presence of human functional sperm.

Conclusion: Our in vivo and in vitro results confirmed that culture of testicular cell suspension and selection of spermatogonial cells could be effective ways for purification and enrichment of the functional human spermatogonial cells. The epigenetic patterns showed that the specific methylation of the evaluated genes at this stage remained constant with no alteration throughout the entire culture systems over time.

Keywords: Azoospermia, Genetic and Epigenetic, Spermatogonial Stem Cells

Introduction

Male infertility is a disorder with complex and multifactorial etiology that caused researchers have been trying to achieve in vitro spermatogenesis (IVS) for a century (1). Epigenetic and genetic modifications roles are essential in spermatogenesis and embryogenesis of clinical approaches (2). In vitro culture and transplantation are two techniques of spermatogonial stem cells (SSCs) resuscitation after cryopreservation that could have a risk to change genetics and epigenetics (3). Consequently, SSCs grown in vitro due to exposure to growth factors and maturation processes can have a higher risk of becoming genetically modified. Therefore, special attention must be paid to the status of in vitro culture and post-transplantation (4).

A low number of SSCs about 0.03% of all germ cells in the rodents and no specific markers for identifying them have hampered rapid success in scientific development (5). Two major developments in SSCs culture include the establishment of the spermatogonial transplantation technique and the identification of glial cell derived neurotrophic factor (GDNF) as a key growth factor for the proliferation of SSCs in vitro (6).

Researchers have used different techniques for the proliferation of SSCs isolated from testis of azoospermic men, such as using human Sertoli cells as a monolayer in the absence of exogenous growth factors (7). Two-dimensional culture systems (2D) resulted in the incomplete proliferation and differentiation stages of SSCs. Thus, three-dimensional (3D) cultures have been
introduced very recently and have been hypothesized to be able to mimic seminiferous epithelium developing male germ cells better (3). Therefore, due to the improvement of culture conditions, one of the most widely used methods in tissue engineering is nanofibers. Poly(lactic acid) (PLA) is a form of an organic polymer obtained from lactic acid dissolved in water and carbon dioxide. Its most important properties are mechanical tensile strength, biocompatibility, and biodegradability. It contributes to mesenchymal stem cells and has chemical and mechanical properties similar to the extracellular matrix (8). In this study, the effect of culturing spermatogonia stem cells with a suspension of testicular cells, Sertoli cells, and culture on PLA nanofiber coated with laminin will be investigated.

Various techniques have been proposed for the isolation of very pure human SSCs (9, 10). Purification is suggested by fluorescence-activated cell sorting (FACS) or magnetic-activated cell sorting (MACS) (11). Panda et al. (12) used Ficoll gradient centrifugation chased by MACS and Thy1 surface marker for extraction of SSCs from growing L. rohita testis. Nevertheless, since there are enormous differences in spermatogenesis details between rodents and mammalians, further studies are essential, especially on humans.

In this study, for the first time, the genetic and epigenetic data on the effect of different in vivo and in vitro conditions evaluated. We aimed to examine genetic and epigenetic changes of human SSCs isolated by MACS and GFrα-1 marker on their proliferation and purification capacity in various cultural systems. SSCs of azoospermic men who had obstructive azoospermia (OA) were cultured in five experimental groups; as a control group, with healthy Sertoli cells of fertile men, with Sertoli cells from men with OA, 3D culture system by PLA nanofiber and suspension groups evaluated. Finally, the SSCs function of the selected culture system after xenograft was considered to evaluate the effect of the length and nature of the culture system on the methylation pattern by the Methylation-specific polymerase chain reaction (PCR) (MSP) method.

Material and Methods

Sample collection

In this experimental study, human research specimens for four experimental groups were taken from men with OA via the intra-cytoplasmic sperm injection (ICSI) program from Shayanmehr Clinic (Tehran, Iran), whose remaining tissue was used for this study. Testicular samples from a fertile man who had an orchidectomy for reasons other than testicular problems were used to obtain healthy Sertoli cells. Testicular samples and the experimental procedure were authorized by Tarbiat Modares University’s National Research Council guidelines (Tehran, Iran). The study goals were clarified to the contributors, and informed consent was taken from patients willing to take part in the research (52/12037). Inclusion criteria for a patient to enter the research are FSH levels between 15-1mIU/ml, testicular volume 6-15 ml, dynamic biopsy with motile sperm, and pathology assessment is positive for spermatogenic cells (13).

Spermatogonial stem cells isolation and in vitro culture

Ten azoospermic men testicular biopsy samples for each experimental group were used and the biopsies specimens were transferred to the laboratory in a medium, within 60 minutes. They were broken into small pieces and placed in Dulbecco’s Modified Eagle medium (DMEM, Gibco, Paisley, UK), got in touch with 14 mm NaHCO₃, (Sigma, St Louis, MO, USA), non-essential amino acids, 100 IU/mL penicillin, and 100 µg/mL streptomycin. The broken pieces of the testis were placed in DMEM, which included 0.5 mg/mL collagenase, 0.5 mg/mL trypsin, 0.5 mg/mL hyaluronidase, and 0.05 mg/mL DNase, for 30 minutes, at 37°C. The gradient of gravity caused the spermatogenic tubules and cells to sediment. After three DMEM washes and the elimination of most interstitial cells, the next digestion step (45 minutes at 37°C) was done in by adding fresh enzymes and media to the fragments of the seminiferous tubule. With centrifugation at 1500 rpm for 4 minutes at 37°C, the cells were separated from the remaining tubule fragments.

Finally, the suspension of testicular cells was incubated in DMEM with FCS 10% and cultured overnight at 37°C and 5% CO₂. Sertoli cells stuck to the bottom of the container faster. In this way, after this period, the top cell suspension containing more germ cells and SSCs was collected.

In the first stage, a total of 2×10⁶ cells were cultured per 12-well plate in five groups for two weeks. You can see the results in our previous study (13). After enrichment of SSCs to enough count, suitable conditions were provided for SSCs purification with MACS. Then we cultured SSCs with and without MACS isolation for one week. Therefore, the cells of each experimental group were divided into two groups part was cultured as before for another week, and part was purified by MACS:

1. Control group, SSCs cultured in the culture dish.
2. Culture of SSCs with men’s own Sertoli cells (Sertoli cells of men with OA). A testicular biopsied specimen was cultured at 37°C after digestion of the second enzymatic period. After 24 hours, the supernatant was removed and the bottom of the dish containing Sertoli cells was cultured with men’s own sorted SSCs.
3. Culture of SSCs with normal Sertoli cells. To provide Sertoli cells for this group, a testicular sample from a fertile man who had an orchidectomy for reasons other than testicular issues was used to achieve healthy Sertoli cells. A testicular specimen was cultured at 37°C after digestion of the second enzymatic period. After 24 hours, the supernatant was removed and the bottom of the dish containing Sertoli cells was cultured with SSCs (13).
4. Culture of SSCs on PLA nanofiber, covered with Laminin.
5. The suspension of biopsied testicular cells from biopsy cultured without separation of SSCs.

Three repetitions of each experimental group were cultured in 34-StemPro for one week with its complement (InVitrogen), 25 μg/ml human insulin, 100 μg/ml transferrin, 60 μM putrescine, 30 nM sodium selenite, 6 mg/ml –D (+) glucose, 30 μg/ml Pyruvic acid, 1 μ/ml –DL lactic acid, 5 mg/ml bovine serum albumin, 2 μM L-glutamine, -25×10⁻⁵ Mercaptoethanol, MEM soluble vitamins, 10⁴ Ascorbic acid, 10 μg/ml -d biotin, 30 ng/ml beta-estradiol, 60 ng/ml bovine serum albumin, 20 ng/ml human epidermal growth factor, 10 μg/ml epidermal growth factor, 10 μg/ml basic fibroblast growth factor (bFGF), 5% FCS, 100 IU/ml human leukemia inhibitory factor (LIF), 10 ng/ml factor class of glial cell and Humanities (GDNF), 10 ng/ml human growth factors, 10 ng/ml derived neurotrophic hormones, 10% FBS, 6 mg/ml transferrin, 60 μM putrescine, 30 nM sodium selenite, 5% D (±) glucose, 30 μg/ml Pyruvic acid, 1 μ/ml –DL lactic acid, 5 mg/ml bovine serum albumin, 2 μM L-glutamine, 25×10⁻⁵ Mercaptoethanol, MEM soluble vitamins, 10⁴ Ascorbic acid, 10 μg/ml -d biotin, 30 ng/ml beta-estradiol, 60 ng/ml Progesterone, 20 ng/ml human epidermal growth factor, 10 ng/ml derived neurotrophic factors, 10 μg/ml basic fibroblast growth factor (bFGF), 5% FCS, 100 IU/ml penicillin, 100 μg/ml Streptomycin (Sigma), and the cells incubated at 37°C with 5% CO₂. We changed the cell culture medium every two days.

Purification of spermatogonial stem cells by MACS

To purify cultured cells, the cells in the plates were washed once with MACS buffer and then an appropriate volume of MACS buffer (based on the plate’s level, number of cells, and the manufacturer’s guideline) was added, and cells were mechanically isolated from the base by a cell scraper and centrifuged. After cell counting, cells were cultured in two groups with and without purification. In the second group, 200 μL buffer was added for all 2×10⁶ cells, and antibodies against GFRα-1 (SC-10716) diluted in 1:50 were added to the cells. Cells were refrigerated for 15 minutes. The column type was chosen based on the number of cells. The column was placed in a separator magnetic field. After washing the column and diluting buffer cells, secondary antibodies attached to Microbead (Milteny Biotec) diluted in 1:10 were added and incubated for half an hour at 4°C on the shaker. After washing with buffer twice and centrifuge, cells were labeled and poured into columns with buffer. Negative cells crossed the column while the GFRα-1 positive cells remained attached.

PLA nanofiber and Laminin preparation

The PLA fibers were made sterile by sinking in ethanol 70% for 2 hours or ultraviolet (14) radiation. Then, 20 μg/mL laminin (Sigma-Aldrich, USA) was poured on them and incubated at 37°C for 2 hours until one night. Before use, it was rinsed with phosphate buffer solution (PBS) and prepared for cell culture. Spermatogonial cell suspension prepared after one-night incubation of the second enzyme lysis was used in this group. The cell load was as follows: first, the cells were suspended in 30 μl of culture medium and a concentration of 3.5×10⁶ cells/mL. The fiber was added to the fiber dropwise and put in an incubator. After two hours, the rest of the culture medium was added. The cell medium was changed every other day.

Spermatogonial and sertoli cells confirmation

The Sertoli cells and SSCs were evaluated for vimentin and GFRα1 markers by immunocytochemistry. The cells were treated with an anti-vimentin antibody. The cells were fixed with 4% formaldehyde and became permeable by 0.2% Triton X100 and clogging with 10% goat serum (Vector, Burlingame, CA) for 30 minutes. The utilized primary antibody (mouse monoclonal anti-vimentin antibody with a dilution, 1:200; Sigma Company, USA) and the rabbit anti-human GFRα1 antibody (dilution 1:100) were added at 4°C and the dishes were incubated 24 hours. The fluorescent-labeled secondary antibody (1:100, Sigma) was added and incubated for 2 hours at 4°C in darkness. The cells were finally mounted with a mounting medium (Vector Laboratories Inc., Burlingame, CA) after three washes with PBS and examined under a fluorescence microscope (IX-71, Olympus).

Quantitative analysis of gene expression

Total RNA was isolated from SSCs derived from all groups, using an RNX-Plus TM (Cinnagen, Iran). RNA concentrations were evaluated by a UV spectrophotometer (Eppendorf, Germany). RevertAidTM first-strand cDNA synthesis kit (Fermentase) with oligo dT primer was used for reverse-transcription of treated RNA. Oligonucleotide PCR primers specific for alpha-6-integrin, beta-1 integrin, PLZF, C-MYC, NANOG, OCT-4, and TBP (internal control) genes were adapted from other primers and synthesized by GenFanAvaran Company.

The Thermal Cycler used SYBR Green and PCR master mix (Cinnagen) for PCR reactions (Applied Biosystems, StepOne TM, USA). Cycling conditions were initiated with a melting period at 95°C for 5 minutes, chased by 40 cycles of melting 30 seconds at 95°C, annealing 30 seconds at 58-60°C and extending 30 seconds at 72°C. Melt curve analysis was performed, and the standard curve for each gene was prepared using serial cDNA dilution from the testis to determine the output. The same run amplified the target gene and the reference gene. The ratio of gene expression was determined using the comparative cycle threshold (CT) method (n=3).

Epigenetic assessment

DNA extraction of spermatogonial stem cells

DNAs of the SSCs in all groups were extracted using a DNA extraction kit (Roche Co) based on the suggested guideline at the end of the culture. The cultured cells were isolated by trypsin and suspended after rinsing in 200 mL PBS. Then, 200 mL binding buffer and 40 μL K proteinase were added and incubated at 70°C. Then, 100 μL isopropanol was added, then centrifuged after being transformed into a filtered tube. Finally, 50 μL elution buffer was added after centrifuge.

DNA methylation by SSS1 enzyme

SSS1 methylase enzyme (Biolabs Co, New England) was used to guide the methylate primers to DNA, according to the instructions. After treatment with sodium...
methylated primers were used for PCR. Enzyme stock 32 mmol was converted to 1600 µmol and incubated at 37°C for 1.5 hours. Heating up to 65°C for 20 minutes would stop the reaction. Then, methylated DNA was treated with SBS, before methylation-specific PCR (MSP) with M primer. Finally, methylated DNA was used as a positive control in MSP with methylated M primer.

**Methylation-specific PCR**

MSP was done by M primer with methylated DNA-modified sequence with SBS, and U primer with non-methylated DNA-modified sequence with SBS. Amplification with M primer showed methylation in CpG zones inside primer sequences, amplification with U primer showed no methylation, and amplification with both primers showed partial methylation in CpG zones inside primer sequences. In the present research, MSP with methylated and unmethylated primers was performed on Integrin α6, Integrin β1, PLZF, C-MYC, and OCT-4.

**Spermatogonial stem cells labeling and transplantation**

To confirm spermatogonial cell identity and function in spermatogenesis, SSCs resulting from cultures were transplanted to twelve NMRI mice, aged 6-8 weeks, with a mean weight of 25 g, kept in separate cages in fair conditions. Initially, the innate spermatogenesis was stopped by an intra-peritoneal injection of 40 mg/kg busulfan. After 4 weeks, the mice were azoospermic. For developing the azoosperma model, 40 mg/kg (Sigma, USA) of Busulfan was injected into each mouse for 4 weeks (15).

To detect and monitor the transplanted cells, spermatogonial colonies were mechanically extracted from culture plates under a reverse microscope. Then, after rinsing with PBS, they were exposed to Dil dye (2 µg Dil per 1 mL of PBS) for 5 minutes at room temperature. Next, they were kept at 4°C in darkness for 20 minutes. After confirming the cell dye under a fluorescent microscope, and three times rinsing in a culture medium, they were transplanted to the recipient mice. The technique used was similar to Brinster’s technique (16). Before transplantation, the recipient mice anesthesia was performed through intraperitoneal injection of 10% ketamine and 20% xylazine. Then, the prepared SSCs were diluted at 10^6 cells per testis in a 10 µL culture medium (17) and micro-injected by 30 Gauge under stereomicroscope guide (Olympus SZ1145, Japan) at a concentration of 10^6/10 µL DMEM to rete testis and finally to the left seminiferous tubule of the mice. To ensure the entrance of the cells to rete testis, 10% trypan-blue dye was simultaneously injected which makes the path visible to the naked eye. The right testis was chosen as the control. After 4, 8, and 16 weeks, the transplanted recipient animals were sacrificed and their testes were separated for assessments under a fluorescence microscope. The cell showing Dil staining was considered transplanted cells. The right testis is considered as the control group without transplantation of donor cells. Also, right side pictures are as phase contrast.

**Assessment of the mouse testes’ weight and epididymis sperms**

The weight of testes was measured by an accurate digital scale after 16 weeks of transplantation in azoospermic mice (n=3). After separating the epididymis of both sides, they were placed in phosphate buffer solution with a pH of 7.4 which had been normalized in the incubator previously. They were then cut for the sperm to exit the epididymis. The phosphate buffer solution containing epididymis parts was incubated at 37°C and 5% CO₂ pressure for 30-45 minutes to increase the capacitated nature of the sperms. Finally, the number of sperms was counted by a hemocytometer under a light microscope and compared between the transplanted and control side.

**Histological studies**

For this part, samples were fixed in Bouin’s fixative solution (Sigma-Aldrich, USA), dewatered by 70-100% alcohol, and elucidated by xylene twice. Then, the samples were dipped with paraffin twice for 1.5 hours, sectioned by microtome with a type C fixed blade (Leitz, Germany), and stained with hematoxylin-eosin (Merck, Germany). The technique was performed in accordance with the manufacturer’s instructions (n=3 per condition). A total of 50 seminiferous tubes were chosen and the number of germ cells (spermatogonia, spermatocyte, and spermatid), as well as the number of cells per surface, were counted in 15 random fields with ×400 magnification under a light microscope.

**Human CatSper expression gene in transplanted mice**

To confirm the presence of human SSCs, quantitative PCR was used to detect human CatSper gene expression in cDNA of the transplanted testis of the mice after 16 weeks (n=3).

**Statistical analysis**

The statistical software SPSS version 16.0 for windows (SPSS Inc., Chicago, IL) was used and graphs were prepared by Microsoft Excel software version 2010. To analyze the data of real-time PCR, first, the raw data were converted to reportable data through available formulas and then analyzed by One-way ANOVA. P values of 0.05 are considered significant.

**Results**

**Confirmation of spermatogonial cells**

The immunohistochemical staining of the isolated Sertoli cells obtained from the seminiferous tubules of testes biopsies of azoospermic men who had OA contained mostly two different cell types: First Sertoli cells which proliferated and formed a monolayer of cells as a feeder layer defined by vimentin. It was observed in the cytoplasm of the Sertoli cells (Fig.1A) around the nucleus (Fig.1B) and merged (Fig.1C). The second type with a spherical outline and two or three exocentric nuclei were spermatogonial cells creating colonies after proliferation (Fig.1D). GFRα-1, which is a spermatogonial stem cell nuclear marker, was found in the obtained colonies (n=3, Fig.1E).
Fig. 1: Spermatogonial and Sertoli cells confirmation. A-C. The immunohistochemical staining of the isolated Sertoli cells obtained from the human testes with detected Vimentin. D. Human spermatogonial stem cell colony. E. Immunofluorescent staining of spermatogonial stem cells (SSCs), detected GFRα1 positive under an immune fluorescence microscope (scale bar: 200 µm).

Isolation of spermatogonial stem cells by MACS

After two phases of crossing the column (repeat separation), Figure 2 shows the immunohistochemistry of expression of GFRα-1 before and after cell isolation with MACS (Fig. 2A, a-d). The percentage of GFRα-1-positive cells by the second isolation phase fraction of MACS was significantly higher than the percentage of positive cells after one isolation phase and before isolation (respectively, 69.01 ± 3.54% to 58.14 ± 2.26% and 37.7 ± 1.53, P<0.05, Fig. 2B).

Results of quantitative polymerase chain reaction

Integrin α6, Integrin β1, and PLZF gene

Without MACS isolation, Integrin-α6, Integrin β1, and PLZF genes expression were significantly higher in testicular suspension cells than in other groups during culture (P<0.05). After cell isolation, the highest Integrin α6, Integrin β1, and PLZF expressions were observed in testicular suspension cells (P<0.05). Also, a comparison of cells isolated with and without MACS showed significantly higher expression of these genes in the isolated groups (P<0.05, Fig. 3A).

NANOG, C-MYC, and OCT-4 gene expression

The expression of the C-MYC gene was lowest in the testicular suspension cells group without isolation of MACS than in healthy Sertoli, simple culture, and nanofiber after one week (P<0.05), but this difference was not statistically significant from men’s own Sertoli cells group (P>0.05). After MACS isolation, it was significantly lowest in testicular suspension cells (P<0.05). Also, a comparison of cells isolated with and without MACS showed lower expression in the isolated groups (P<0.05).

The expression of Nanog after one week, without isolation of MACS, was lowest in testicular suspension and highest in simple culture (P<0.05), but this difference was not statistically significant between the men’s own Sertoli cells group and nanofiber groups (P>0.05). After isolation with MACS, the expression of NANOG was lowest in testicular suspension and highest in the control group (P>0.05). The gene expression was lower in isolated than in non-isolated groups.

OCT-4 gene expression was highest in the simple culture, with and without cell isolation (P<0.05). The comparison of cells isolated with and without MACS showed no difference in gene expression (P>0.05, Fig. 3).

Epigenetic results of MSP in different groups during culture

MSP results with methylated primer for Integrin α6, β1, and PLZF gene in all cultured cells had a similar pattern in all culture systems and remained non-methylated. The size of the proliferation fragment for Integrin α6 for methylated primers was 100 bp and for non-methylated primers 101 bp. They were 203, and 205 bp for Integrin β1 gene, while for PLZF gene they were 125, and 130 bp, respectively. Methylation pattern did not change in C-MYC and OCT-4 gene during culture and it remained in partial methylation. The size of the proliferation fragment was 140 bp for methylated and non-methylated primers in PLZF gene and 105 bp for OCT-4 (Fig. 4).
Comparing Spermatogonial Cells In Vitro Xenotransplant In Vivo

Fig.3: Quantitative gene expression analysis by qRT-PCR. A. Integrin α6, B. β1, C. PLZF, D. NANOG, E. C-MYC, and F. OCT-4, during spermatogonial cells culture in the studied groups. In each group, the expression level of a gene in each sample is normalized to TBP, as an internal control. The level of expression of each sample is also calibrated to a calibrator (the cells derived from second enzymatic digestion). α; Significant differences with other groups each time (P≤0.05), β; Significant differences between MACS+ and MACS- groups (n=3, P<0.05, P≤0.05).

The results of in vivo assessment

Regarding the culture results of the previous steps as well as epigenetic studies, the suspension culture group was considered the best group, and the resulting cells were selected for transplantation. The azoospermic status of the mice before transplantation was confirmed (Fig.5A).

Monitoring the transplanted cells

The results of the sections’ assessment showed that after 4 weeks, the cells were placed at the base of seminiferous tubes (Fig.5B); after 8 weeks, the tracked cells were shown in the diameter of seminiferous tubes (Fig.5C); and after 16 weeks, some seminiferous tubes of the transplanted testis (Fig.5D, left side) contained spermatozoa. Also, endogenous spermatogenesis was observed in the control testis (right side, Fig.5B-D).

Comparison of the testes’ weight and number of sperms in epididymis between the transplanted and control testis

The mean testis weight of the azoospermic SSCs transplanted and control mice showed a significantly higher mean weight than azoospermic (sham group: without SSCs transplantation) and the opposite side testis of the transplanted azoospermic mice that were not transplanted after 16 weeks (0.08 ± 0.00058, 0.095 ± 0.0015, 0.057 ± 0.0032, 0.063 ± 0.0051g respectively, P<0.05). The mean testis weight of the transplanted group was significantly lower than the mean testis weight of normal healthy adult mice (P<0.05, Fig.6A).

Assessment of the number of sperms in epididymis revealed that the transplanted and control testis had significantly higher mean sperms than the sham and opposite side testis groups after 16 weeks (26 ± 5, 31 ± 9, 18.7 ± 4, 16.3 ± 5 respectively, P<0.05). The mean sperm count of the transplanted group was significantly higher than the mean sperm count of the sham and opposite side testis groups (P<0.05). The number of sperms in the control testis of the transplanted mouse was similar to the transplanted group (P>0.05, Fig.6B).

Histopathology of testis sections in the transplanted and control testis

Based on Figure 6C-E, the mean number of spermatogonia, spermatoocytes, and spermatids was higher in the transplanted testis than in the control and sham (P<0.05), but it was
significantly lower than that of normal healthy adult mice (P<0.05). The mean number of cells in the control testis of the transplanted mouse was similar to that of the sham group (P>0.05).

Assessment of CatSper gene in the transplanted testis

CatSper gene was evaluated after 16 weeks of transplantation in the azoospermic mouse model. The expression of this gene in the transplanted testis showed proliferation and presence of the human CatSper gene in the transplanted testis. Nevertheless, the results showed significantly lower CatSper gene expression in the transplanted testis than in the human testis (P<0.05). The expression of this gene was zero in the control testis (Fig.6F).

Discussion

Optimization of a system for the proliferation and differentiation of male germ cells is a valuable tool for managing male infertility and spermatogenesis regulation (18). One of the important regulators in different spermatogenesis processes is epigenetic modifications (19). Genetic and epigenetic structures of chromatin are essential for fertile sperm production (20). Hence, we used MACS for human SSCs purification by GFR-α1 positive marker. GFRα1 is a self-renewal-related or pre-meiotic gene, expressed in undifferentiated SSCs such as A single, A paired and spermatogonia A allele (21). The MACS isolated SSCs were cultured in different culture systems and compared for genetic and epigenetic expression between five experimental groups; i. Control two dimensional culture, ii. Co-culture of SSCs with
Sertoli cells of men with OA, iii. Co-culture of SSCs with healthy Sertoli cells, iv. Culture of SSCs on PLA nanofiber, and v. Culture of testicular cell suspension. Researchers have similarly investigated the isolation of human SSCs through the FACS method (9, 22). Meanwhile, the high costs and time-consuming nature of FACS may limit its application in cell isolation. Thus, we used two-step enzyme lysis and incubation of the resulting cells for one night and isolation of suspended spermatogonial cells on the next day, as required in different groups. The results of the present study confirmed GFR-α1 (GDNF receptor) as an effective marker in improving SSCs isolation, which has been previously suggested by Godmann et al. (23) as well. It has been confirmed that GDNF supplies the necessary items for the growth and maintenance of human SSCs in the medium and generally shows the suitability of MACS with GFR-α1 for isolation and enrichment of human SSCs (24). Miltenyi et al. (25) suggesting MACS as a fast and simple separation system for large immunologic cells. Baert et al. (26) cultured MACS-enriched epithelial cells in the interstitial cell-laden scaffolds (CD49f+/CLS). They observed double-cell compartment testicular constructs. Cell spheres showed in the pores after cell seeding on CFS and CLS. The elongated spermatids were observed in 66% of TC/CFS. Differentiation was achieved in all and 33% of CD49f+/CLS constructs, respectively.

The MACS enriched and non-enriched (SSCs cultured without MACS sorting) SSCs were cultured in the control and treated groups; co-culture of SSCs with Sertoli cells of men with OA, co-culture of SSCs with healthy Sertoli cells, the culture of SSCs on PLA nanofiber, and culture of testicular cell suspension. We observed significantly the highest expression of Integrin α6 and PLZF genes in the testicular suspension cells group and lowest expression in the simple culture group than in other groups. Integrin β1 gene expression was highest in testicular suspension cells and lowest in the PLA nanofiber group, which confirmed that testicular suspension cells could effectively purify and enrich the functional human spermatogonial cells. Integrin α6 and Integrin β1 are premeiotic markers and they have a connection to laminin and collagen proteins in the base membrane of seminiferous tubes (6). Nevertheless, the expression of OCT-4 gene was not significantly different among groups. The gene expression of OCT-4 and PLZF is related to GDNF and affects the self-renewal of SSCs (27, 28). The three-dimensional culture, with the aid of an extracellular matrix, enables cells to organize properly and imitate the spermatogonial epithelium (29). The use of three-dimensional culture on PLA nanofiber with a laminin layer in the present study showed the superiority of this method over simple culture, regarding germ-cell specific gene expression. We used PLA, because of their biodegradability and biocompatibility, Which is quickly made by electrospinning and creates a three-dimensional non-woven grid. Eslahi et al. (8) cultured SSCs seeded on PLLA with the control groups and suggested that PLLA increases the colony formation of human SSCs in the culture system. Also in the present study, a superior co-culture with Sertoli cells in comparison with the control group was observed. It might be SSCs culture with GDNF-secreting Sertoli cells acting as spermatogonial cell renewal regulator. Koruji et al. (30) co-cultured human SSCs and Sertoli cells and observed an increase in the count and diameter of SSC colonies. The gene expression of pluripotency genes showed minimum expression of C-MYC gene and Nanog in testicular suspension cells, especially after isolation with MACS with increased gene expression during culture in all isolated groups. Rajpert-De Meyts et al. (31) could isolate multivalent germ cells from the adult human testis by using suitable culture conditions to isolate embryonic stem cells. They observed that these cells express OCT-4 proteins but do not express NANOG.

According to our previous study, the suspension culture group was regarded as most similar to testes’ micro-environment (13). We observed that the other culture groups, such as Sertoli and PLA nanofiber, which kept the nature of human SSCs also confirms the importance of the presence of extracellular matrix, micro-environment, and their signaling. On the other hand, germ cells often show genetic and epigenetic changes in vitro, and SSCs seem to maintain relative genetic stability. SSCs characteristics did not change and cells were not differentiated suggesting the stability of this culture technique, which could be due to the protective effect of GDNF against differentiation (32). It can be a reason for prolonged epigenetic changes after culture among different groups with and without MACS. This may limit the assessment of gene expression and DNA methylation to only some printed genes. Because of this, comparative genomic hybridization is not able to detect small genetic changes. Goossens et al. (33) evaluated the DNA methylation pattern in a paternally methylated gene (Igf2), a maternally methylated gene (Peg1), and a non-imprinted gene (α-Actin). The spermatids obtained from the 3D-I system have similarities in global gene profile and DNA methylation compared to in vivo spermatids. They used MACS for human spermatogenesis in-vitro, by isolating GPR125+ spermatogonia from the testes of OA patients.

Transplantation of SSCs is a fertility restoration option that has already been introduced as a convenient method in animals (16). Studies on the imprinting situation after SSCs transplantation are limited but show that implantation does not change. The results of the present study on transplantation of human SSCs, cultured with healthy Sertoli cells to seminiferous tubes of a recipient mouse, showed that it can result in spermatogenesis with a donor origin. Although transplantation of human SSCs is not likely to produce sperm in mice, it may activate endogenous spermatogenesis-stimulating factors. Thus, we evaluated sperm production in mice after 16 weeks. When the SSCs from the donor are transplanted to the seminiferous tubes of an infertile recipient, the germ cells of the donor migrate to the lateral base of the tubes in the seminiferous tubes of the recipient, then proliferate, produce new colonies, and start spermatogenesis.
with a donor origin (35). Although initial studies have experimented with mice models (16), this technique can be useful in spermatogenesis studies of other animals as well. Indeed, some studies have reported successful heterografts from mouse testis to hamsters that resulted in spermatogenesis of mouse and hamster (36). There are several studies that transplanted human SSCs into the mouse testis and reported that SSCs adhere to the seminiferous tubules after 2 weeks (22, 34). Mohaqiq et al. (34) isolated human SSCs and confirmed them by PLZF protein. They transplanted SSCs to adult azospermia mouse testes and studied them after two weeks. The results revealed that the number of SSCs was significantly more than those in the control group. IHC studies and qRT-PCR indicated that the PLZF was only expressed in the transplantation groups. The results of SSCs transplantation in our study were similar to others (9, 34).

GCT, as the ultimate goal of these cellular studies, could successfully restore spermatogenesis in animal models and resolve infertility, which is considered the gold standard (37). Further research continues to elucidate different aspects of GCT for successful experiments on humans. Despite the scientific development in stem cells, human SSCs culture is still a controversial issue. The results of the present study can dynamically add to the knowledge of researchers and clinicians and is an important step toward future clinical use for male infertility, which was the strongest strength of the present study. In addition, we could successfully achieve an appropriate number of cells through a two-phase culture which was an important limitation in previous studies (38). Yet, the current study had some limitations, including the fact that the transplantation experimented on mice and thus, the results cannot easily be generalized to humans. Further research on mammals can add to the results of the present study.

Conclusion

The epigenetic pattern showed that the specific methylation of the evaluated genes at this stage remained constant throughout the entire culture system over time and the culture conditions did not alter the methylation pattern. Also, MACS could increase the efficiency of human SSCs isolation and purification by 69% with the testicular suspension group showing the highest expression of germ cell genes (Integrin α6, β1, and PLZF), and lowest gene expression of C-MYC gene and NANOG, among the tested groups. Further, the proposed culture systems could maintain the cell-specific genetic and epigenetic contain and the suspension cells group known as the best system for SSCs culture in vitro. Thus, the results indicate the ability of purification and proliferation of functional cells in the suspension culture.

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Authors’ Contribution

M.Z; Performed the experiments, data acquisition, data analysis and interpretation, and drafting the manuscript. M.M.; Was the conductor of the study, participated in study design, edited the manuscript, also participated in the finalization of the manuscript, and also approved the final draft. S.J.M., M.N.; Advised for genetic and epigenetic assessment, real-time PCR technique, extracted mRNA and produced cDNA. M.R.N; Advised sample collection and separated SSC from samples. M.K., F.A.; Performed transplantation and participated in statistical analysis, and edited the manuscript. All authors were involved in the drafting and revision of the draft manuscript.

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