INTRODUCTION
The pluripotent properties of stem cells have attracted considerable interest in both basic research and clinical applications.1-4 There are three main types of stem cells, namely embryonic stem (ES) cells, adult stem cells, and the induced-pluripotent stem (iPS) cells. However, ethical issues as well as tumorigenicity and immune rejection preclude the applications of ES cells in clinic.4-6 Similarly, the iPS cells encounter several severe hurdles, e.g. viral transduction, low efficiency, and tumor-causing risk,7 although they can be induced to generate male germ cells.8 Due to these obvious drawbacks, scientists are devoting to seeking alternative sources of pluripotent stem cells from adult tissues. Notably, spermatogonial stem cells (SSCs) might become an ideal candidacy because of their unique and important plasticity, including self-renewal, differentiation, dedifferentiation, and transdifferentiation.7,8,10 SSCs are a small subpopulation of type A spermatogonia in mammalian testis where they can self-renew and differentiate to produce sperm in the course of normal development throughout life. We have recently shown that SSCs from cryptorchid patients could be induced to differentiate into round spermatids with high efficiency, and tumor-causing risk,11 Strikingly, numerous studies by peer and us have recently demonstrated that SSCs can directly differentiate into morphologic, phenotypic, and functional cells of other lineages. Direct conversion to the cells of other tissues has important significance for regenerative medicine. SSCs from azoospermia patients could be induced to differentiate into spermatids with fertilization and developmental potentials. As such, SSCs could have significant applications in both reproductive and regenerative medicine due to their unique and great potentials. In this review, we address the important plasticity of SSCs, with focuses on their self-renewal, differentiation, dedifferentiation, transdifferentiation, and translational medicine studies.

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SELF-RENEWAL AND DIFFERENTIATION OF SPERMATOGONIAL STEM CELLS
Spermatogonial stem cells undergo self-renewal to maintain the stem cell pool while they differentiate into sperm in mammalian testis.5,18 Spermatogonia are classified into three types, including type A, intermediate, and type B cells, based upon their morphological characteristics.19 Type A spermatogonia can be subdivided into A<sub>app</sub> (A<sub>0</sub>), A<sub>pure</sub> (A<sub>1</sub>) and A<sub>aligned</sub> (A<sub>2</sub>) cells.20,21 SSCs are very rare, with only approximately 0.02%-0.03% of all germ cells in adult testis;22 nevertheless, thousands of spermatids are produced each day, reflecting an unlimited potential of SSC division and differentiation. We illustrated the self-renewal and differentiation of the rodent and human SSCs in Figure 1. In rodents, A<sub>0</sub> spermatogonia can divide symmetrically to produce new A<sub>0</sub> stem cells or they divide asymmetrically to give rise to A<sub>0</sub> spermatogonia and A<sub>1</sub> spermatogonia that are characterized by an intercellular cytoplasmic bridge. The A<sub>0</sub> spermatogonia continue to divide and form chains of A<sub>0</sub> spermatogonia (4, 8, 16, and 32 cells) that are able to differentiate into type A<sub>1</sub> to A<sub>4</sub> intermediate (In) and B spermatogonia. Type B spermatogonia further differentiate to primary spermatocytes, secondary spermatocytes, and eventually spermatids via two meiotic divisions. There are distinct cell type and biochemical phenotypes between rodent and human male germ line stem (GS) cells.
Additionally, FGF2 is required for the survival and differentiation of SSCs in vitro.

We have revealed that GDNF activates CREB/ATF-1 family members and c-fos transcription via the Ras/ERK1/2 pathway to promote DNA synthesis and proliferation of SSCs. Therefore, BMP4 and SCF have been employed to probe the differentiation of SSCs into pachytene spermatocytes.

It has been reported that SCF and RA could induce spermatogenesis from cryopreserved neonatal SSCs and spermatogonial cell lines to differentiate into spermatocytes and spermatids in vitro, and this process is c-KIT-dependent.

For example, the receptor RARα of RA can increase SCF and BMP4 expression, whereas it reduces GDNF secretion. However, currently little is known about how SSCs balance the self-renewal and differentiation. To unveil more signaling molecules and pathways or targets regulating SSC self-renewal and differentiation would contribute to a better understanding the fundamental principles of stem cell biology and treatment of male infertility.

The microenvironment or the niche precisely regulates the fate decisions of SSCs. There are differences in the roles of signaling pathways in controlling SSCs between species. For instance, JAK/STAT signaling pathway has been demonstrated to stimulate the self-renewal and maintenance of Drosophila SSCs. In contrast, STAT3 signaling pathway is shown to be required for the differentiation of mouse SSCs. Sertoli cells, a key component of the niche, produce certain growth factors, e.g. glial cell line-derived neurotrophic factor (GDNF), fibroblast growth factor 2 (FGF2), bone morphogenetic protein 4 (BMP4) and stem cell factor (SCF), that play important roles for the self-renewal and/or differentiation of SSCs. GDNF has been demonstrated to be essential for the survival and proliferation of SSCs in vivo and in vitro. We have revealed that GDNF activates CREB/ATT-1 family members and c-fos transcription via the Ras/ERK1/2 pathway to promote DNA synthesis and proliferation of SSCs. Mouse SSCs proliferate over a 5-month period when cultured with basic fibroblast growth factor (bFGF) (also known as FGF2), GDNF, leukemia inhibitory factor (LIF), and epidermal growth factor (EGF), and both bFGF and GDNF stimulate Ras pathway via Src family kinases. Additionally, FGF2 is required for the survival and proliferation of mouse SSCs. On the other hand, BMP4 and SCF have been shown to promote the differentiation of SSCs. Although BMP4 is unimportant for the survival of undifferentiated spermatogonia, it can induce SSC differentiation through changing cell adhesion properties and activation of Smad1/5/8 pathway by up-regulation of c-KIT expression. Deletions or mutations of SCF or its receptor c-KIT result in sterility in mice due to lack of differentiating spermatogonia, implicating that SCF/c-KIT pathway is essential for SSC differentiation. SCF also contributes to the proliferation of primary spermatagonia in culture, and the division of type A and A spermatogonia is c-KIT-dependent.

To induce the differentiation of SSCs and their precursors with an objective to generate male germ cells in vitro, as we summarized in Table 1, might provide male gametes for treating male infertility. Early in 1910s, scientists started to examine spermatogenesis process using testicular tissue culture, and male germ cells could enter meiotic prophase in vitro. Notably, pachytene primary spermatocytes were observed after 11 days of culture from neonatal mouse testes. Gas-liquid interphase organ culture system was developed and it was optimized with culture parameters, e.g. temperature, incubation atmosphere, pH, nutrients (pyruvate, vitamin A, E, and C), for the differentiation of rat and human spermatogonia from testis tissues into pachytene spermatocytes. Cell culture and coculture methods were employed to probe the differentiation of SSCs in vitro. It has been reported that preleptotene spermatocytes progress toward later stages of meiotic prophase in association with Sertoli cells that provide necessary microenvironment in vitro. Immortalized mouse type A spermatogonial cell line was induced to generate haploid spermatids in vitro, however, whether these spermatids have function of fertilization remains unknown. Several groups have reported that rodent type A spermatogonia and early male germ cells could differentiate into haploid cells when co-cultured with Sertoli cells. However, gene expression patterns of spermatids derived from the type A spermatogonia were abnormal and no offspring was obtained by micro-insemination. Furthermore, the fertilization potential of those round spermatids was not tested, and the efficiency was very low. Human spermatogonia derived from nonobstructive azoospermic patients were coaxied to differentiate into haploid cells. Although the efficiency of differentiation was low, the round spermatid-like cells possessed normal chromosome status and could activate human oocytes after injection into the cytoplasm. Recently, a breakthrough has been reported that SSCs from freshly and cryopreserved neonatal mouse testes could produce spermatids and sperm with serum-free culture media. It is worth noting that this sperm could give rise to healthy and reproductively competent offspring. We have recently induced SSCs from cryptorchid patients to differentiate into haploid spermatids with fertilization and developmental potentials, which...
would provide an invaluable source of autologous male gametes for treating male infertility of azoospermia patients. Since gene expression patterns of human SSCs are different after short- and long-term culture, it is of great significance to develop efficient approaches for maintaining self-renewal and differentiation of SSCs.

**DEDifferentiation of Spematogonial Stem Cells to Embryonic Stem-like Cells**

Spematogonial stem cells have previously been considered unipotent because they are only committed to produce sperm in the course of normal development. However, several lines of evidence suggest that SSCs can acquire pluripotency. First, teratomas which contain derivatives of all three germ layers occur exclusively in the gonads, and ES-like cells derived from SSCs in culture with certain growth factor could form teratomas; second, primordial germ cells (PGCs) which differentiate into gonocyte and subsequently into SSCs can produce embryonic germ cells that are similar to ES cells in potency. Pluripotency of germ cells can be maintained postnatal. Therefore, investigators propose that SSCs are pluripotent. Great efforts from a number of pioneering studies have proven this notion. Lastly and more importantly, numerous groups have demonstrated that SSCs both from mouse and human testis could be induced to become ES-like cells that differentiated into derivatives of three embryonic germ layers (Table 2). In 2004, pluripotent ES-like cells were first generated from SSCs of neonatal mice by Dr. Shinohara and colleagues when they cultured SSCs in GS cells medium with addition of GDNF, bFGF, EGF, and LIF. Notably, these ES-like cells could differentiate into various types of cells, e.g., hematopoietic cells, vascular cells, and spontaneously beating myocytes. Interestingly, more neural lineage cells and heart muscle cell colonies from ES-like cells were found than from ES cells. Typical teratomas and chimerism were seen in ES-like cells derived from SSCs and there were no significant histological differences from teratomas derived from ES cells. These findings strongly implicate that neonatal mouse SSCs can acquire pluripotency. ES-like cells derived from adult mouse testis were also successfully achieved. Their phenotypic characteristics and gene expression profiles were similar to ES-like cells induced by Dr. Shinohara group. ES-like cells derived

| Table 1: The differentiation of SSCs and other male germ cells in vitro |
|----------------|--------------------|------------------|---------------------|
| **Species** | **Approaches** | **Major findings** | **References** |
| Newborn mice | Watch-glass method on the surface of a clot: equal parts of fowl plasma and fowl embryo extract | Gonocytes→pachytene spermatocytes | 52 |
| Adult men | Gas-liquid interphase method: EMEM, pyruvate, NEAA, glutamine, FCS | Preleptotene→pachytene spermatocytes | 55 |
| Adult men | Tissue culture: Parker 199 supplement fructose, coconut milk, FBS, FSH, LH, testosterone | Preleptotene spermatocytes→telophase II spermatocytes | 56 |
| Prepubertal rats | Co-culture: EMEM, pyruvate, glutamine, NEAA, FBS, FSH, testosterone, transferrin, EGF, retinol | Preleptotene spermatocytes→pachytene spermatocytes | 59 |
| Mouse SSC lines | Cell culture: DMEM, NEA, sodium pyruvate, glutamine, FCS | Spermatogenesis→round spermatids | 60 |
| Mouse SSC line | Cell culture: DMEM+SCF | Type A spermatogonia→spermatocytes | 33 |
| Adult men | Co-culture: Vero cell condition medium, FSH and testosterone | Type A spermatogonia→spermatocytes | 61 |
| Neonatal rats | Co-culture: DMEM, FBS, glutamine, pyruvate, NEAA, EGF, IF, FSH, insulin, transferring, retinol, retinal, testosterone, dithyrotetosterone, nucleoides solution, human growth hormone | Type A spermatogonia→round spermatids | 62 |
| Neonatal mouse | Organ culture: aMEM, FBS | Type A spermatogonia→functional sperm | 64 |
| Azospermic patients | Co-culture: Ko-DMEM, KSR, FSH, GDNF, testosterone, RA | Type A spermatogonia→haploid cells | 63 |

SSC: spermatogonial stem cell; NEAA: nonessential amino acids; EMEM: Eagle’s minimum essential medium; FCS: fetal calf serum; LH: luteinizing hormone; FBS: fasting blood sugar; FSH: follicle stimulating hormone; RA: retinoic acid; GDNF: glial cell line-derived neurotrophic factor; KSR: knockout serum replacement; EGF: epithelial growth factor; DMEM: Dulbecco’s modified Eagle’s medium; SCF: stem cell factor; IFG: insulin-like growth factor; aMEM: a-minimal essential medium

| Table 2: The dedifferentiation and transdifferentiation of SSCs |
|----------------|----------------|----------------|---------------------|
| **Species** | **Starting cells** | **Induction protocol** | **Pluripotency** | **References** |
| Neonatal mice | Early type of spermatogonia | GS medium+GDNF+EGF+bFGF+LIF | Derivatives of three germ layers, teratoma formation, EBs and chimaeras | 10 |
| Adult mice | STRA8+ cells | Basic medium+GDNF | Derivatives of three germ layers, teratoma and EBs | 12 |
| Adult mice | GPR125+ germine progenitors | GS medium+GDNF | Derivatives of three germ layers, teratoma formation, EBs and chimaeras | 17 |
| Human | CD49f+ cells with collagen and laminin selection | Basic medium+LIF and/or GDNF | Derivatives of three germ layers and teratomas | 13 |
| Human | Testicular cells | ES medium on MEF | Derivatives of three germ layers, EBs, no large teratoma | 70 |
| Human | Testicular cells | DMEM+bFGF+TGF-β | Derivatives of three germ layers, no large teratoma | 71 |
| Human | Testicular cells | GS medium+GDNF+EGF+bFGF+LIF | Derivatives of three germ layers, no teratoma | 14 |
| Adult mice | Testicular cells | SSCs mixed with mammary epithelial cells in mammary fat pads | Functional mammary epithelial cells | 74 |
| Neonatal mice | Early type of spermatogonia | Combined with inductive mesenchymes in vivo | Tissues of three germ layers, e.g., prostatic, uterine, and skin epithelium | 15 |
| Neonatal mouse | SSCs | Step 1: Nodal+Wnt+bFGF | Functional hepatocytes | 16 |

GS medium: the culture medium published by Kanatsu-Shinohara et al.
Basic medium: DMEM: 15% FCS; 1% NEAA; 1% L-glutamine, and β-mercaptoethanol; GDNF: glial cell line-derived neurotrophic factor; EGF: epithelial growth factor; LIF: leukemia inhibitory factor; GS: germ line stem; bFGF: basic fibroblast growth factor; MEF: mouse embryonic fibroblast; EBs: embryoid bodies; SSC: spermatogonial stem cell; ES: embryonic stem; DMEM: Dulbecco’s modified Eagle’s medium; TGF-β: transforming growth factor beta; EGF: epithelial growth factor; HGF: hepatocyte growth factor; HCM: hepatocyte culture medium; OSM: oncostatin M; NEAA: nonessential amino acids
from adult mice could differ into ectodermal-, mesodermal- and endodermal-derived tissues and they produced mature teratomas and chimaeric mice with live offspring, a gold standard for pluripotency. In 2007, ES-like cells were induced from GPR125+ germine progenitors of adult mice, and they produced cells and tissues of three germ layers and could contribute to chimera embryos. Although different induction protocols were used, ES-like cells were uniformly formed from neonatal and adult mouse testes, reflecting that SSCs become pluripotent ES-like cells. On the other hand, due to no specific and unique marker for SSCs, highly purified SSCs were rather hard to obtain, which pose several questions, such as the origin of ES-like cells and the dedifferentiation mechanism. Kim et al. identified three transitional stages during SSC dedifferentiation to ES-like cells, including SSC stage, intermediate state SSCs, and ES-like morphology stage, which might be helpful for better understanding the mechanisms controlling the dedifferentiation of SSCs to ES-like cells.

A number of groups attempted to obtain autologous ES-like cells from human SSCs. Excitingly, Conrad et al. first demonstrated the successful establishment of pluripotent ES-like cells derived from spermatogonial stem cells of adult human testis. They obtained a highly pure SSC population by using CD49f and followed by sequential matrix selection with collagen and laminin, which has important significance to define the origin of ES-like cells. Human ES-like cells were generated from human SSCs with pluripotent characteristics and expression profiles similar to human ES cells when they were cultured with LIF. Human ES-like cells derived from SSCs could form a teratoma. Derivation of human ES-like cells has also been achieved from a testis biopsy or donor testes. Although these human ES-like cells expressed markers of pluripotency and formed embryoid bodies (EBs), they didn’t produce large teratomas. It is speculated that human ES-like cells may not have been reprogrammed sufficiently to produce teratomas, which might be a great merit to clinical application of SSCs. As such, human SSCs derived ES-like cells can be used to generate various kinds of cells for cell-based therapy and tissue engineering for human diseases. Importantly, the dedifferentiation of SSCs to ES-like cells doesn’t involve virus vectors and ethical issues, which is much safer compared to human iPSCs or human ES cells.

**DIRECT TRANSDIFFERENTIATION OF SPERMATOGLONIAL STEM CELLS TO THE CELLS OF OTHER LINEAGES**

Spermatogonial stem cells arise from PGCs and they share similar expression of certain key genes (e.g. Oct-4 and Nanog) for ES cells. This indicates that the closest equivalent of ES cells in vivo is probably SSCs. Since SSCs can acquire pluripotency to become ES-like cells that subsequently differentiate into other lineage tissues, it is reasonable to presume that SSCs can directly transdifferentiate into other cell types without the pluripotent status. Boulanger and colleagues direct the transdifferentiation of testicular stem cells into functional mammary epithelial cells. However, the prerequisite is that they must mix spermatogenic cells with dispersed mammary epithelial cells, followed by transplanting them into the mammary fat pad. Engraftment of SSCs alone cannot form mammary epithelium, which suggests that SSCs cannot transit to the mammary epithelium de novo. It has been reported that SSCs-enriched cells alone from neonatal mice transdifferentiate directly into tissues of all three germ layers, including prostatic, uterine, and skin epithelium. Furthermore, the organs and tissues transdifferentiated from SSCs expressed molecular, histological, and functional markers of the appropriate epithelium. Engraftment of SSCs can promote recovery in a rat Parkinson’s disease model, and rat SSCs transdifferentiate to functional dopaminergic neuron-like cells via mesenchymal–epithelial interactions. Importantly, we have recently demonstrated that mouse SSCs are able to transdifferentiate directly into morphological, phenotypic, and functional hepatocyte-like cells when they are cultured with several growth factors in vitro. SSCs first converted into hepatic stem-like cells which subsequently differentiate into small hepatocytes and mature hepatocyte-like cells. Notably, we did not detect the gene expression for ES cells in the whole transdifferentiation process, such as SSEA-1, SSEA-4, Nanog and TRA-1-81, which suggests that no ES-like cells was formed during the transdifferentiation process. Very recently, we have induced the transdifferentiation of mouse SSCs into functional dopaminergic neurons in vitro (unpublished data).

As illustrated in Figure 2, SSCs can give rise to a wide range of other type cells directly, which implies that they have important significance in regenerative medicine. First of all, the direct transdifferentiation of SSCs to mature and functional cells without the process of de-differentiation to ES-like cells and EB formation could simplify the reprogramming procedure of cells. Secondly, the conversion of SSCs using growth factors without gene modification could be much safer to generate mature cells for cell therapy and tissue engineering for human disease. Nevertheless, there are several issues to be defined prior to the application of cells derived from SSCs in clinic. First, the mechanisms of direct transdifferentiation are poorly understood. One concept is that there is a distinct subpopulation of pluripotent SSCs that can directly transdifferentiate into cells of another lineage. Izadyar et al. have identified two distinct populations in the GS cells, and only the POU5F1+/c-KIT+ cells are pluripotent. Another viewpoint is that all SSCs are capable of becoming pluripotent once removed from their niche. It is possible that the pluripotency of SSCs is repressed by the seminiferous tubular microenvironment in which they reside. Once outside of this niche, they recover original characteristics and convert to another lineage depending on the particular niche in which they are placed. Third, the necessary prerequisite to clinical application is to obtaining sufficient numbers of highly purified SSCs
from human testis. Therefore, it would be crucial to identify specific and unique SSC markers. We have isolated human SSCs using 2-step enzymatic digestion and magnetic-activated cell sorting, and notably, the isolated SSCs can be cultured in an undifferentiated state for 2 weeks. Nevertheless, culture condition needs to be optimized for long-term survival and expansion of human SSCs. Finally, it is important to explore the optimal induction protocols with higher transdifferentiation efficiency of SSCs to mature and functional cells. Enough transition cells from a limited number of SSCs are necessary for cell-based transplantation therapy. Therefore, extensive studies are required to eliminate those hurdles.

PERSPECTIVES AND CONCLUDING REMARKS

As addressed above, SSCs represent a novel and significant cell source for both reproductive and regenerative medicine, due to their unlimited potentials including pluripotency, self-renewal, differentiation, and transdifferentiation. Generation of mature and functional cells from SSCs have certain advantages over ES cells and iPSCs, including no ethical concern, lower frequency of tumorigenesis, and no immune rejection. On the other hand, certain issues await to be clarified, e.g. the dedifferentiation and transdifferentiation mechanisms, optimal induction protocols, and origin of the ES-like cells. Due to no specific and unique marker available for SSCs, the true cellular origin of ES-like cells remains controversial. Given the rapid progress in SSC research, it is no doubt that SSCs would be eventually utilized from the bench to bedside in the near future.

AUTHOR CONTRIBUTIONS

ZH designed the outline of the manuscript. ZC, ZL and ZH wrote the manuscript, and ZH finalized and approved the manuscript.

COMPETING INTERESTS

All authors declare no competing interests.

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