Spermatogonial stem cell enrichment using simple grafting of testis and in vitro cultivation

Jung Jin Lim1, Dong Won Seol2, Kyung Hee Choi1, Dong Hyuk Shin1, Hyung Joon Kim1, Seung-Hun Song3 & Dong Ryul Lee1,2

1Fertility Center of CHA Gangnam Medical Center, College of Medicine, CHA University, Seoul, 135-913, Korea, 2Department of Biomedical Science, College of Life Science, CHA University, Seoul, 135-081, Korea, 3Department of Urology, CHA Gangnam Medical Center, College of Medicine, CHA University, Seoul, 135-913, Korea.

Enrichment of spermatogonial stem cells (SSCs) from the mammalian adult testis faces several limitations owing to their relatively low numbers among many types of advanced germ cells and somatic cells. The aim of the present study was to improve the isolation efficiency of SSCs using a simple tissue grafting method to eliminate the existing advanced germ cells. Sliced testis parenchyma obtained from adult ICR or EGFP-expressing transgenic mice were grafted heterotropically under the dorsal skin of nude mice. The most advanced germ cells disappeared in the grafted tissues after 2–4 weeks. Grafted tissues were dissociated enzymatically and plated in culture dishes. During in vitro culture, significantly more SSCs were obtained from the grafted testes than from non-grafted controls, and the isolated SSCs had proliferative potential and were successfully maintained. Additionally, EGFP-expressing SSCs derived from graft parenchyma were transplanted into busulfan-treated recipient mice testes. Finally, we obtained EGFP-expressing pups after in vitro fertilization using spermatozoa derived from transplanted SSCs. These results suggest that subcutaneous grafting of testis parenchyma and the subsequent culture methods provide a simple and efficient isolation method to enrich for SSCs in adult testis without specific cell sorting methods and may be useful tools for clinical applications.

Spermatogenesis is the process by which undifferentiated germ cells divide and mature, resulting in the sustainment of male fertility via the daily production of millions of spermatozoa in the testis. The foundation of this process lies in spermatogonial stem cells (SSCs), which undergo self-renewal and produce daughter cells by undergoing complicated differentiation processes1–4. In the past few decades, techniques for long-term, in vitro maintenance of SSCs have been greatly improved by co-culturing on feeder cells and/or in defined medium containing various growth factors, such as glial cell line-derived neurotrophic factor (GDNF), glial cell line-derived neurotrophic factor receptor-α1 (GFR-α1), basic fibroblast growth factor (bFGF), epidermal growth factor (EGF), and leukemia inhibitory factor (LIF)5–7. Similarly, other researchers have reported that the selection of testicular cells expressing integrin α6 (CD49f), integrin β1, CD9, Thy-1 or GFR-α1 resulted in significantly increased efficiency in SSC colonization8–11. These studies indicated that a relatively high ratio of SSCs might be required for the efficient isolation of cell lines with biological potential. Indeed, the testes of neonatal or transgenic animals that are amenable to various cell sorting methods have usually been used as starting materials for the establishment of SSC lines12,13. Meanwhile, because of the small ratio of SSCs in the human adult testis14,15, conventional methods for isolating adult derived-human SSCs may limit clinical trials16,17. Recently, some groups have applied various methods to develop an enrichment procedure for adult-derived SSCs. For example, SSCs can bind to laminin but not to fibronectin or collagen type IV18. When laminin-binding (lamB) and laminin-non-binding (lamNB) GFP+ germ cell populations selected from 19-day-old mice were transferred to the testes of recipient males, only the lamB fraction was substantially colonized. This finding suggested that SSCs were greatly enriched in the lamB fraction, which represented approximately 5% of the total germ cell population. However, unlike the pre- or peri-pubertal testis, this protocol was not applicable to the isolation of SSCs from adult testes. Other researchers have taken advantage of the sensitivity of advanced germ cells to high core body temperature. When the testis of most mammals is retained in the body cavity, spermatogenesis fails to occur. Additionally, if the mature testis is surgically fixed in the abdomen, the mature stages of germ cells are lost. This condition is known as cryptorchidism. An experimental procedure for cryptorchidism...
has been used in a variety of laboratory studies\textsuperscript{19,20}, but this procedure is not appropriate for human clinical trials. In the third model, a testis tissue graft in an immunodeficient host mouse has been shown to model the structural integrity of the testicular tissue; this model therefore facilitated the accessibility and controlled manipulation of testicular function\textsuperscript{11,12}. However, early studies using this procedure were focused on the completion of spermatogenesis \textit{in vivo} following long-term transplantation (6 ~ 12 months) of immature testis\textsuperscript{21,22} and not on the isolation or maintenance of SSCs.

We hypothesized that short-term (1 ~ 4 weeks), ectopic adult testis tissue grafting would lead to a new method for isolating or maintaining SSCs and could serve as a model system that would be applicable in human clinical trials via the simple grafting of the patient’s own tissue to eliminate the advanced germ cells present in the adult testis. To verify this possibility, in the present study, SSCs were isolated and cultured from grafting tissues using a collagen type IV-coated dish and then evaluated their self-renewal capacity. The biological potential of the SSCs was confirmed by the production of advanced germ cells and offspring after transplantation to busulfan-treated, infertile recipient mice.

**Results**

Evaluation of grrafted testicular parenchyma survival and morphology. Fig. 1A shows the typical morphology of the tissue sections of non-grafts and whole or slice grafts. At the time of grafting, the seminiferous tubules included all types of cells: Sertoli cells, SSCs and differentiated germ cells. Two weeks after grafting, 14/20 (70.0%) and 17/20 (85.0%) of the grafted tissues (whole tissue and slice transplantation, respectively) contained germ cells and Sertoli cells in the dorsal skin. However, round spermatids and sperm were not observed in the seminiferous tubules. In the whole tissue and slice groups, 4/20 (20.0%) and 3/20 (15.0%) of the grafted tissues, respectively, were observed to have fully sclerotic tubules. To determine whether SSCs were included in the grafting tubules, we used immunohistochemical staining to detect the SSC markers GFR-\(\alpha\)-1 and Thy-1. The results showed that 14/16 (87.5%) and 14/17 (82.4%) of the whole and slice grafted tissues, respectively, contained SSC marker-positive cells in the dorsal skin. Four weeks after grafting, the numbers of sclerotic tubules increased to 7/10 (70%) and 6/10 (60%) in the whole tissue and slice groups, respectively. Interestingly, no seminiferous tubules exhibited GFR-\(\alpha\)-1/Thy-1-positive staining (Table 1). Therefore, the 2-week grafting groups were selected for subsequent experiments.

To analyze the viability of cells within the seminiferous tubules, twelve slides, four from each of the three groups, were stained using TUNEL methods. In the non-grafts (control), the percentage of apoptotic cells (TMR-red stained) was 4.76 \(\pm\) 4.97%. Although the total number of cells (DAPI-stained) within the seminiferous tubules was significantly decreased compared with the control, the percentages of apoptotic cells were not different between the whole or slice graft tissues (8.89 \(\pm\) 8.22% vs.9.48 \(\pm\) 8.83%) (Fig. 1B,C).

**In vitro culture of SSCs derived from grafted testicular parenchyma.** Some of the dissociated cells from the testicular parenchyma (cells unattached to the collagen-coated dish) were cultured in StemPro-34 SFM supplemented with 1% KSR and with four growth factors (LIF, EGF, FGF2, and GDNF). Owing to the gelatin coating of the culture plate, some somatic cells attached and proliferated strongly for AP staining (Fig. 2B). To confirm that an AP staining-positive clump was an SSC clump, we performed co-staining for E-cadherin (a marker for SSCs) and 3\(\beta\)-HSD (a marker for interstitial cells). As shown in Fig. 2C, AP was co-expressed in both the E-cadherin-positive and some of the 3\(\beta\)-HSD-positive cells. However, 3\(\beta\)-HSD was not detected in SSC clumps showing AP- and E-cadherin-double positive staining. Attached somatic cells gradually decreased during the second 7 days of culture, but after passing, they re-formed clumps that subsequently became loosely attached to dishes and continued to proliferate for more than 8 passages (>2 months).

**Localization of SSC markers and characterization of cultured cells.** It is well known that Thy-1 and GFR-\(\alpha\)-1 are expressed on mouse and human SSCs, and this was confirmed by co-staining with E-cadherin (a marker for SSCs) in seminiferous tubules (Fig. 3A and 3B). Thus, we hypothesized that the ratios of Thy-1 and GFR-\(\alpha\)-1-positive cells could be increased after dissolution of differentiated germ cells using the grafting method. Immunohistochemical analysis of normal adult mouse testis showed that Thy-1 and GFR-\(\alpha\)-1-positive cells (putative SSCs) were detected on the basement membrane, but their ratios were low. However, in the grafted testicular tissues, although their tubules appeared small in size, the ratio of cells showing Thy-1 and GFR-\(\alpha\)-1 signals were relatively higher than those in fresh tissue (Fig. 3A and 3A'). Observed under high magnification, as expected, Thy-1 and GFR-\(\alpha\)-1 were strongly expressed on the surface membrane of cultured SSCs. During \textit{in vitro} culture, the colonization efficiency of the SSCs was higher in the grafted testis than in non-grafted controls (Fig. 3C and 3C').

**Flow cytometrical characterization and RT-PCR of cultured cells.** After 2 weeks of culture, the relative ratios of the SSCs were determined using FACS with Thy-1 and GFR-\(\alpha\)-1. The ratios of CY3 (Thy-1) cells in testicular cells that were obtained from whole and slice grafted testes were 85.00% and 83.40%, respectively, but the ratio from non-grafted testis was 7.33%. Similarly, the ratios of FITC-labeled (GFR-\(\alpha\)-1) cells in testicular cells that were obtained from whole and slice grafted testes were 78.92% and 68.26%, respectively, but the ratio of those cells from non-grafted testis was 3.93% (Fig. 4A).

Additionally, to characterize the cultured cells that were derived from the non-grafted, whole and sliced testis grafted groups, we analyzed the expression of spermatogenesis-specific genes by RT-PCR. First, we analyzed the expression of Gfr-\(\alpha\)-1, lin28, and ngn3, which are specifically expressed in undifferentiated spermatogonia. Cultured cells from the whole grafted group demonstrated high expression of the mRNAs of Gfr-\(\alpha\)-1 and lin28 compared with cells from the non-grafted and slice grafted groups. Interestingly, the expression of ngn3 was not different in any of the three groups. However, stra8 and piwi2, which are specifically expressed in differentiated spermatogonia and spermatocytes, were highly expressed in cultured cells of the non-grafted group and were weakly expressed in both grafted groups. However, the expression of c-kit was similar across the groups. Additionally, the TH2B and TP-1 genes, which are specifically expressed in spermatocytes and spermatids, were decreased in cells from the grafted groups. Mvh, which is expressed in all stages of germ cell development, was slightly decreased in cells of the grafted groups, but not significantly. These results indicated that the majority of cultured cells included SSCs but did not include advanced-stage germ cells, such as spermatocytes and spermatids (Fig. 4B).

**Generation of functional spermatogenic cells by transplantation of cultured cells into the testis of busulfan-treated males.** To determine the function of SSCs that had been isolated \textit{in vivo} and proliferated \textit{in vitro}, we next transplanted them into the testis of infertile recipient mice. For these experiments, SSCs isolated from EGFP-expressing (C57BL/6-Tg (CAG-EGFP)) transgenic mice were used. The efficiency of isolating SSCs from the transgenic mice was similar to that of the previous experiment using wild type mice.
Following a culture period of 4 weeks, isolated SSCs were harvested and transplanted into the seminiferous tubules of busulfan-treated adult mice via an efferent duct. Some EGFP-positive germ cells in recipient testes were observed under UV light or in paraffin sections 4 weeks after transplantation (Fig. 4B).

To produce embryos, spermatozoa or elongated spermatids were collected from the recipient testes 4–8 weeks after transplantation and injected into BDF1 oocytes. After ICSI, 80/101 (79.2%) eggs survived and 71/80 (88.7%) developed to the 2-cell stage within 24 hours in culture. In some preimplantation embryos produced by ICSI, a strong EGFP signal (6/10) was detected at the blastocyst stage. However, some cultured blastocysts did not express EGFP (4/10), which seems to have been due to the production of embryos by fertilization from the recipient’s own sperm that had re-initiated

### Table 1 | The presence of germ cells in the testicular parenchyma of ICR mice after grafting into dorsal skin of nude mice

| Transplanted type | Transplanted period | No. of parenchyma | No. of atrophic parenchyma (%)* | No. of parenchyma showing expression of SSC-markers (%) |
|-------------------|---------------------|-------------------|--------------------------------|-----------------------------------------------|
| Whole testis      | 2 weeks             | 20                | 4 (20.0)                       | 14/16 (87.5)                                 |
|                   | 4 weeks             | 10                | 7 (70.0)                       | 0/3 [0.0]                                    |
| Sliced testis     | 2 weeks             | 20                | 3 (15.0)                       | 14/17 (82.4)                                 |
|                   | 4 weeks             | 10                | 6 (60.0)                       | 0/4 [0.0]                                    |

*Whole parenchyma that had fully degenerated seminiferous tubules.
spermatogenesis after busulfan treatment (Fig. 4C). To generate offspring, the embryos (30 blastocysts) were transferred into the oviducts of two pseudopregnant females, which resulted in the delivery of 7 live pups (four EGFP-expressing pups and three non-EGFP-expressing pups, Supplementary Table 1 and Fig. 4D). As expected, genotyping analysis by PCR revealed that among the seven pups, four expressed EGFP (Supplementary Fig. 1). These results indicated that SSCs isolated in vivo can be differentiated into functionally advanced germ cells and are capable of producing normal offspring.

Discussion
Although successful isolation and proliferation of SSCs has recently been reported\(^1-10,13,20-27\), these methods are still limited because high efficiency is only accomplished in SSC-enriched, prepubertal animals.
or by cell sorting-based strategies using specific antibodies. However, for clinical purposes, the isolation of SSCs from adult testes is more important than isolation from prepubertal testes. Moreover, a high ratio of advanced germ cells in the adult testis may reduce the isolation efficiency of cell sorting using specific antibodies. Therefore, the development of a novel, efficient method for the enrichment and isolation of SSCs from adult testis is a prerequisite for the treatment of male infertility and the study of spermatogenesis in vitro. Many studies have focused on germ cell production of SSCs, and several recent studies have reported that SSCs can be converted to pluripotent stem cells in specific culture conditions. However, unlike neonatal testicular cells, the conversion or isolation of pluripotent stem cells from adult testis has rarely been reported in mammalian and human systems. Thus, the difficulty of isolating sufficient numbers of SSCs has likely hindered the progression of their study and application.

Figure 3 | Immunolocalization of E-cadherin, GFR-α1 and Thy-1 in adult mouse testis after a 2-week grafting and in their cultured SSCs. (A) Cross-sections of seminiferous tubules in an adult mouse testis stained for Thy-1 and GFR-α1. Thy-1 and GFR-α1, which indicate an SSC-positive signal, are shown in red and green (Thy-1; CY3 and GFR-α1; FITC) and were found in cells located at the basement membrane. (A’) Graphical representation of the results of the graft group. The ratios of SSC-marker-positive cells were significantly increased when compared with the non-graft group. (B) Cross-sections of seminiferous tubules in an adult mouse testis stained for GFR-α1 and E-cadherin. GFR-α1 and E-cadherin, which indicate an SSC-positive signal, are shown in green and red (GFR-α1; FITC and GFR-α1; CY3) and were found in cells located at the basement membrane. The right panel shows a high magnification image of the rectangular area. (C) Cultured SSC colonies from graft tissues at 2 weeks were fixed and stained with Thy-1 and GFR-α1. Thy-1 and GFR-α1 were abundantly expressed on the membrane and in some of the cytoplasm of the SSC colonies. C’. During culture (60 days), SSCs proliferated continuously and maintained similar size morphology. Note: Mock primary Ab/DAPI: DAPI counter-stained SSCs without a primary antibody. The data are presented as the mean ± SEM. a vs b: significantly different (P < 0.05). Scale bars, 20 μm.
Figure 4 | Flow-cytometric analysis and RT-PCR of cultured SSCs from adult mouse testis tissues 2 weeks after grafting into recipient nude mice. (A) The percentage of Thy-1 and GFR-α1-positive cells in the non-graft group was only 7.33 and 3.93%, respectively. The percentage of Thy-1 and GFR-α1-positive cells in the graft (whole) group was 85.00 and 78.92%, respectively, which was significantly higher than that in the non-graft group. The percentage of Thy-1 and GFR-α1-positive cells in the graft (slice) group was 83.40 and 68.26%, respectively, which was also significantly higher than that in the non-graft. (B) Germ cell-specific mRNA expression patterns by real time RT-PCR. Gfr-α1, lin28 and ngn3 mRNA were used as undifferentiated spermatogonia marker genes. c-kit, piwll2 and stra8 were used for differentiating spermatogonia and spermatocytes. TH2B was used for spermatocytes. TP-1 mRNA was used as a spermatid marker gene, MVh was used for all germ cells, and 18S ribosomal RNA was used as a control gene.
To enrich SSCs from the mammalian adult testis, the experimental cryptorchid procedure was applied and resulted in the death of most of the germ cells, leaving only undifferentiated spermatogonia 2 months after surgery in mice, which led to a 5- to 25-fold enrichment of SSCs. McLean et al. reported that the colonization (an SSC property) of germ-line stem cells from hyperthermia-exposed testes was 6- to 19-fold higher than in controls when calculated based on the number of colonies and the area of colonization.

Grafting of testis tissue was developed as a tool for androgen substitution in the 1950s and has been subsequently applied to the study of steroidogenesis and spermatogenesis. Rathi et al. and Schoenfeld et al. reported that testicular stem cells continuously proliferated in the aging testis after grafting. In the present study, we introduced a testis tissue grafting method for the enrichment of SSCs from adult testis connected with the simultaneous cultivation of these cells because tissue grafting is potentially be a better strategy for germ cell preservation and transplantation in humans than the cryptorchid or hyperthermia-exposed methods. In fact, as a model for application, testis grafting into a recipient mouse enriched the SSC population, and culture using collagen selection further increased the concentration of SSCs by decreasing the number of advanced germ cells and somatic cells (Fig. 2). Additionally, isolated SSCs proliferated continuously in vitro culture conditions for 2–3 months, and they kept their stem-cell characteristics. To confirm that the cultured cells were SSC colonies, we performed an immunocytochemical analysis to detect GFR-α1 and Thy-1 expression in the cultured germ cell clumps. As expected, GFR-α1 and Thy-1 were strongly expressed in the cultured SSCs (Fig. 3 and Fig. 4). The purity of the SSC population was gradually decreased (Fig. 4B). Additionally, these observations explain why the specific sorting procedure for germ cells could be skipped in our graft/culture system using one type of ECM (collagen type IV).

The technique for the transplantation of cultured cells into the seminiferous tubules of recipient males is well established in mice. In the present study, SSCs were isolated from the testes of EGFP-expressing transgenic mice and propagated (Fig. 5A), and harvested cell suspensions were microinjected into seminiferous tubules via rete testis. Transplanted SSCs colonized the basement membrane of the seminiferous tubules and begin proliferating (Fig. 5B). After 4–6 weeks, differentiated germ cells begin to fill the tubule from the basement membrane toward the lumen, and a few spermatids or spermatozoa appeared in the lumen. Using the ICSI technique, the donor-derived, differentiated germ cells produced embryos and offspring (Fig. 5C and 5D), suggesting that the biological potential of SSCs isolated by our grafting/culture method can be well maintained.

Recent advances in chemotherapy and radiotherapy have significantly improved the remission and complete recovery rates of cancer patients. However, because germ cells are highly susceptible to cytotoxic treatments, iatrogenic loss of fertility has emerged as a major side effect of successful treatment. In males, sperm freezing is an established method to preserve germine cells and is performed routinely in clinics for patients who wish to preserve their fertility before undergoing treatment for malignancy. Unfortunately, the technique cannot be applied to prepubertal patients, who do not have sperm but have different types of germ cells. Therefore, the tissue grafting germ cell culture method may provide a new avenue for SSC-based cell therapy for the preservation of an individual male’s SSCs and re-introduction of those cells into the testis after...
patient recovery while avoiding re-introduction of cancer cells during tissue transplantation. Additionally, this technology may increase the efficiency of studies on SSCs for germ cell and multipotent stem cell production from adult male patients.

Methods

Animals. All mice were housed under a 12 hr/12 hr light/dark cycle in a temperature- and humidity-controlled room. Donor testses for transplantation were obtained from 6- to 8-week-old adult, outbred (wild-type) C57BL/6 mice (C57BL/6-Tg(CAG-EGFP)) transgenic mice (SLC, Inc., Hamamatsu, Japan), Six- to 8-week-old adult BALB/c-Nu (Samtaco Co., Ltd.) male mice were used as the testis recipients.

Adult testis transplantation. Donor adult mice were killed by cervical dislocation, and their testes were collected and/or cut transversally into 3–5 slices. The tissues were maintained in ice-cold buffered medium (Dublcco’s modified Eagle’s medium), with added antibiotics and 10% fetal bovine serum (FBS; Invitrogen) at room temperature for up to 30 min until the recipient was surgically prepared for grafting. Recipient nude mice were anesthetized using ketamine (80 mg/kg body weight) and xylazine (10 mg/kg body weight) through the right carotid incision. The somatic scrotal skin was ligated, and the testicles were removed. The scrotal skin was then closed using sutures, and two dorsal scrotal incisions were made in the skin and body wall. Donor testes or testicular tissues were placed under the dorsal skin using fine forceps and closed by sutures. The mice were maintained in groups of four per cage with food and water available ad libitum for 2–4 weeks.

Isolation of grafted parenchyma and in vitro culture of SSCs. The testes parenchyma were harvested at 2 weeks (n = 20) and 4 weeks (n = 10) after grafting. The dorsal skin of each sacrificed mouse was removed, the grafts were excised, and 1/4 of the tissue was fixed in Bouin’s solution overnight at room temperature for histological and immunohistochemical analyses. The remaining graft parenchyma were dissociated enzymatically in buffer containing 5 mg/ml collagenase (Type IV; Sigma-Aldrich, St Louis, MO), 10 μg/ml DNase I and 1 mg/ml hyaluronidase (Sigma-Aldrich) for 5 min with agitation at 37°C. Dissociated cell suspensions were then plated and incubated on collagen-coated dishes in selective medium composed of Dublcco’s modified Eagle’s medium (DME; Invitrogen, Carlsbad, CA) containing 20% FBS, 10 μM 2-mercaptoethanol (Invitrogen) and 1% non-essential amino acids (Invitrogen) at 37°C under humidified conditions of 5% CO2 in air. Over the next 12–24 h, unattached cells were harvested and plated on 0.1% gelatin-coated dishes (1–1.5 × 107 cells/cm²) in serum-free germline stem cell (GS) medium with modified supplements, as previously described. The composition of the GS medium was as follows: StemPro-34 SFM (Invitrogen) supplemented with 6 mg/ml D(G)-glucose, 5 × 10⁻⁵ M β-mercaptoethanol, 1 μM D(-)lactic acid, 2 mM L-glutamine, 30 μM pyruvic acid, 10⁻² M ascorbic acid, 60 ng/ml progesterone, 30 ng/ml β-estradiol (Sigma-Aldrich), 0.2% BSA (ICN Chemicals, Costa Mesa, CA), 100 U/ml penicillin, 100 μg/ml streptomycin, 1x Insulin-Transferrin-Selenium supplement, 1x MEM vitamin solution, 1x MEM non-essential amino acids, 20 ng/ml mouse EGF (Invitrogen), 10 ng/ml human bFGF (Invitrogen), 1x Knockout™ serum replacement (KSR, Invitrogen), 10 ng/ml rat GDNF (R&D systems, Inc., Minneapolis, MN) and 10 μM USGRO (Chemicon, Temecula, CA). During culture, some remaining somatic testicular cells attached to the bottom of the dish. Adult testicular cell-derived SSCs proliferated slowly and then formed clump-like structures on the somatic cells 2 to 4 weeks after seeding. Cultured cells were dissociated by trypsin treatment and re-plated every 7 to 10 days. Cultured cells were maintained at 37°C in 5% CO2, and the medium was changed every 2 days.

Histological and immunocytochemical characterization of graft tissue and cultured SSCs. After fixation, tissue samples were embedded in paraffin and cut into 5- to 7-μm-thick sections. Each section was stained with hematoxylin and eosin for morphological evaluation. For the detection of apoptotic cells, sections were permeabilized by incubation in 0.5% Triton X-100 for 1 h at room temperature, and a TUNEL assay was performed using an In Situ Cell Death Detection Kit as described by the manufacturer (Roche Diagnostics, Indianapolis, IN). Sections were washed twice with PBS and incubated with the TUNEL reaction mixture (10 μl of terminal deoxynucleotidyl transferase and 90 μl of nucleotide mixture) for 60 min at 37°C in a humidified atmosphere in the dark. To detect alkaline phosphatase (AP) activity, cultured cells were fixed in 4% paraformaldehyde at room temperature for 1 min, washed twice with PBS and stained with an alkaline phosphatase substrate solution (10 ml FRV-Alkaline Solution, 10 ml Naphth AS-BI Alkaline Solution; Alkaline Phosphatase Assay Kit, Sigma-Aldrich) for 30 min at room temperature. AP activity was detected colorimetrically (red) by light microscopy. For the detection of SSC-specific markers, tissues on cell or cell colonies in phosphate-buffered saline (PBS, pH 7.4) for 30 min at 4°C. Slides were then stained with primary antibodies (1:100–500 dilutions) against E-cadherin (Santa Cruz), Thy-1 (BD Biosciences), GFR-91 (Chemogen) or JH-3 (BD Biosciences), and embryo transfer (ET). BDF1 female mice were superovulated by injection of 5 IU of pregnant mare serum gonadotropin (PMSG, Folligon, Intervet Co., Holland) followed 48 hr later by injection of human chorionic gonadotropin (hCG, Chorion, Illinois, USA). For recipients, 14 hr post-hCG, 0.4% NIH-B6D2F1F1 female mice were used. The donor testes were stored in a Petri dish containing modified human tubal fluid medium (HTF, Irvine Scientific, Santa Ana, CA) at room temperature. After being washed, the oviducts were placed in fresh medium, and cumulus-oocyte complexes were released.

Flow cytometric analysis. Flow cytometric analyses were performed using a standard protocol. SSC colonies were dissociated in trypsin-EDTA, and aliquots of 10⁶ cells were suspended in 0.1 ml of PBS containing 2% fetal bovine serum (PBS/FBS) and incubated with primary antibodies. To detect SSC-specific cells, the cells were incubated with 10 μg/ml anti–mouse CD90.2 (1:50; 20) and 4 weeks (n = 20) and third experiments (4 mice) to test their in vivo biological activities. To transfer the cells into the seminiferous tubules of a recipient testis, we used a transfer technique via the efferent ducts. Cultured SSCs were dissociated and suspended in DPBS medium containing 0.4% trypan blue (Sigma-Aldrich). The recipient mice were anesthetized, and the testes were exteriorized through a midline abdominal incision and immobilized. The injection pipette was constructed from a three-inch-length of borosilicate glass with an internal diameter of 0.3–0.5 mm (World Precision Instruments, Inc., WT100-3). The glass was drawn on a Kopf pipette puller (Model 750), which created two potential injection pipettes (40–60 μm outside diameter (o.d.)). The glass pipette tip was inserted into the efferent duct using a dissecting microscope, and the pressure in the injection tubing was raised until the cell suspension flowed into the seminiferous tubule. The flow was monitored based on the color change. A cell suspension of approximately 3–5 μl (less than 1 × 10⁶ cells/ml) was injected into each recipient testis, which filled 50–90% of the tubules in each testis. Four to 8 weeks after transplantation, the recipient testes were collected and analyzed by EGFP expression using a fluorescent microscope, and then collected haploid germ cells were used for embryo production.

Preparation of gametes, intracytoplasmic sperm injection (ICSI), embryo culture and embryo transfer (ET). BDF1 female mice were superovulated by injection of 5 IU of pregnant mare serum gonadotropin (PMSG, Folligon, Intervet Co., Holland) followed 48 hr later by injection of human chorionic gonadotropin (hCG, Chorion, Illinois, USA). For recipients, 14 hr post-hCG, 0.4% NIH-B6D2F1F1 female mice were used. The donor testes were placed in a Petri dish containing modified human tubal fluid medium (HTF, Irvine Scientific, Santa Ana, CA) at room temperature. After being washed, the oviducts were placed in fresh medium, and cumulus-oocyte complexes were released.
from the ampulla using Dumont forceps. In the ICSI procedure, the cumulus cells were dispersed by incubation for 3–5 min in HTF medium containing 0.1% hyaluronidase (Sigma-Aldrich); after being washed, the oocytes were maintained in KSOM medium at 37°C in 5% CO₂ in air until ICSI. Testicular sperm from the recipients were collected in M2 medium by excision with a pair of fine scissors and forceps, followed by squeezing in a 50-μl drop of M2 in mineral oil. The sperm were allowed to disperse at 37°C for 30 min, and each suspension was transferred to a 50-μl Eppendorf tube for centrifugation at 1500 rpm for 5 min. Sperm were mixed with 40 μl of 10% polyvinylpyrrolidone (PVP-360) in M2, followed by placement in a culture dish for microinjection. Injections were performed using a PMM-150 FU piezo-impact unit (Prime Tech) and micromanipulators using a blunt-ended, mercury-containing glass needle (inner volume, 6–7 μl). After 15 min of recovery at room temperature in M2 medium, the surviving oocytes were returned to mineral oil-covered KSOM and cultured at 37°C in an atmosphere of 5% CO₂ in air. For embryo culture, 50 μl of KSOM covered with mineral oil were equilibrated overnight at 37°C in a humidified atmosphere of 5% CO₂. After 92–96 hr of culture, in vitro grown blastocysts were transferred into the uterus of a 2.5-day pseudopregnant female that had been mated with the vasectomized male mice. The newborns were observed using a UV illuminator.

Statistical analysis. Unless otherwise specified, each experiment was carried out using at least three replicates. Data are expressed as the means ± SEM. The statistical significances of the differences between treated groups were evaluated by one-way analysis of variance (ANOVA) using a log-linear model in the Statistical Analysis System (SAS, Cary, NC, USA). P values < 0.05 were considered statistically significant.

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Additional information

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