Amniotic Fluid Stem Cells Prevent Follicle Atresia and Rescue Fertility of Mice with Premature Ovarian Failure Induced by Chemotherapy

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Abstract

Chemotherapy used to treat cancer may cause irreversible premature ovarian failure (POF). Of late, amniotic fluid stem cells (AFSCs) provide a novel source for regenerative medicine because of their primitive stage, low immunogenicity, and easy accessibility. In this study, we isolated AFSCs from transgenic mice that ubiquitously express enhanced green fluorescence protein (EGFP). These AFSCs exhibited morphologies, immunophenotypes, and mesoderm trilineage differentiation potentials similar to mesenchymal stem cells (MSCs). Further, AFSCs proliferated faster than MSCs and expressed OCT4, a marker for pluripotency. To investigate their potential in recovering fertility in POF model, AFSCs were transplanted into the ovaries of mice with POF six weeks post induction using chemotherapeutic drugs, busulfan and cyclophosphamide. AFSCs could rescue the reproductive ability of mice with POF by preventing follicle atresia and sustaining the healthy follicles. Notably, the transplanted AFSCs did not differentiate into granulosa and germline cells in vivo. After one month, the decreased numbers of transplanted AFSCs accompanied with the reduced beneficial effects indicated that the therapeutic efficacy were directly from AFSCs. These findings demonstrated the therapeutic effects of AFSCs and suggested the promise of AFSCs for treating infertility and POF caused by chemotherapy.

Introduction

Chemotherapy (CTx) is commonly used for treating various malignancies and can improve the survival rate for cancer patients [1]. A potential adverse effect of chemotherapy is to decrease the number of ovarian follicles, disrupt the menstrual cycle, induce infertility, and cause irreversible premature ovarian failure (POF) [2–4]. In addition, early onset of menopause can result in the reduction in the quality of life. POF is associated with risks of cardiovascular disease [5], osteoporosis [6], and psychiatric diseases such as depression [7]. A recent study showed that bone marrow transplantation can rescue the fertility of mice with CTx [8]. Subsequently, more studies proved that transplantation of mesenchymal stem cells (MSCs) could restore partial ovarian function in a mouse model of CTx-induced POF [9,10]. These findings suggested the potential of MSCs to treat ovarian failure in humans, but its cellular and molecular mechanisms are still unknown. Furthermore, the proliferative capacity of MSCs in vitro is limited, and its acquisition is highly invasive [11]; therefore, an alternative cell source in clinical POF applications may be preferred.

To address this issue, we reasoned that amniotic fluid stem cells (AFSCs) may serve as an alternative option because they are readily harvested from amniotic fluid, undergo rapid self-renewal, maintain a normal karyotype in long-term culture [12], and tally with all the properties defined in the minimum criteria for MSCs [13–16]. Compared with other MSCs, AFSCs are easy to harvest, have a shorter doubling time and express the embryonic stem cells (ESCs) marker OCT4 [17], whereas unlike ESCs and induced pluripotent stem cells, they do not develop into teratomas in vivo [18]. Further, evidence indicated that transplanted allogeneic AFSCs have low immunogenicity and can escape immune rejection [19]. The potential advantages of AFSCs for treating POF are further suggested by their use in the studies of regenerative medicine to repair damaged tissues or organs, such as muscle [14], kidney [20], lung [21], or heart [22].

In this study, we evaluated the efficacy of AFSCs in a mouse model of POF and investigate the potential mechanism of this therapeutic effect. We established and characterized AFSCs from
transgenic mice ubiquitously expressing enhanced green fluorescence protein (EGFP), which enables us to trace the fate of AFSCs after transplantation. To evaluate whether AFSCs can rescue the female fertility, we transplanted AFSCs expressing EGFP (EGFP-AFSCs) into the ovaries of mice with POF, assessed ovarian function and traced the fate of transplanted AFSCs within the ovaries of recipients.

Materials and Methods

Animals and Ethics Statement

ICR mice aged 4–6 weeks were purchased from the Laboratory Animal Center of Medical College of National Taiwan University (Taipei, Taiwan). EGFP transgenic mice that ubiquitously express EGFP were as previously described [23]. In brief, EGFP-expressing transgenic mice lines were created by pronuclear microinjection of ICR strain zygotes with a construct (pCX-EGFP) that expresses EGFP under control of the β-actin promoter [24–26]. Mice were housed under a 14–10 h light–dark cycle at 25°C±2°C with food and water provided ad libitum. The Institutional Animal Care and Use Committee of National Taiwan University approved the experimental protocol (NTU-102-EL-10).

Isolation and Culture of EGFP-AFSCs

Mice were euthanized by sequential CO2 asphyxiation and cervical dislocation at day 11.5 of pregnancy. The uterus was isolated and 26-gauge needles were used to collect amniotic fluid from each EGFP-expressing conceptus. Amniotic fluid was added to the wells of a 24-well cell culture dish (#39024, TPP, Trasadingen, Switzerland) containing 1 mL of Minimum Essential medium supplemented with 3.7 mg/mL NaHCO3 (#SH30070-03, GE Healthcare Life Sciences, South Logan, Utah), 100 IU/mL penicillin, and 100 mg/mL streptomycin (#S7671, Sigma-Aldrich), 10% (v/v) fetal bovine serum (FBS) (#SH30070-03, GE Healthcare Life Sciences, South Logan, Utah), 100 IU/mL penicillin, and 100 mg/mL streptomycin (#15140, Gibco, Grand Island, NY). Cells were incubated at 37°C in a humidified atmosphere of 5% CO2. Every three days, 0.5 mL of non-adherent cells were removed and replaced with 0.5 mL of fresh culture dishes and incubated as described above. Non-adherent cells were removed by changing the medium every three days. Cells (70% confluent) were removed from the dishes by incubation at 37°C with 0.25% (w/v) trypsin/ethylenediaminetetraacetic acid (EDTA) (#2500056, Gibco) for 5 min.

Flow Cytometry Analysis

To assess the immunophenotypes of cultured AFSCs, we performed a flow cytometric assay. Cells were reacted in the dark for 30 min at 4°C according to the manufacturers’ protocols with the following phycoerythrin (PE)-conjugated monoclonal antibodies: CD11b (#12-0112), CD29 (#12-0291), CD31 (#12-1031), CD34 (#12-1051), CD44 (#12-0441), CD45 (#12-0451), MHC-I (#12-5985), MHC-II (#12-5321) and Sca-1 (#12-3981). A FACSA disappear (BD Biosciences, Franklin Lakes, NJ) was used to analyze 1×106 cells, and the data were processed using the FCS Express software (Version 4.0; Denovo Software, Los Angeles, CA). All antibodies included isotype-matched control antibodies were purchased from eBioscience (San Diego, CA).

Mesoderm Trilineage Differentiation of EGFP-AFSCs

In order to investigate whether the cultured EGFP-AFSCs have the mesoderm trilineage differentiation potentials as MSCs, the differentiation ability test is based on the induction system with modifications. To induce adipo genesis, near-confluent monolayers of EGFP-AFSCs were cultured in MEM-α supplemented with 10% FBS, 10 μg/mL insulin (#I6634), 1 μM dexamethasone (#D4902), 0.5 mM isobutyl-methylxanthine (#I5879), and 100 μM indomethacin (#17378) for 14 days with medium changes every three days. Cells were fixed with 4% paraformaldehyde (PFA) (#P6140) at room temperature for 10 min, and lipid droplets were stained using Oil Red O (#O9755) [27].

To induce osteogenesis, confluent cells were incubated in MEM-α supplemented with 10% FBS, 0.1 μM dexamethasone, 10 mM β-glycerophosphate (#G9091) and 50 μM ascorbic acid (#A8960) for 21 days with medium changes every three days. Bone matrix mineralization was evaluated using Alizarin Red S staining (ARS) (#A533) [27]. A pellet culture system was used to induce chondrogenesis with MEM-α supplemented with 1% FBS, 6.25 μg/mL insulin, 50 μM ascorbic acid and 10 μg/mL transforming growth factor (TGF)-β1 (#240-B-002, R&D Systems, Minneapolis, MN) with twice weekly medium changes. Proteoglycan production was determined using toluidine blue O (#T3280) staining [28]. All chemicals were purchased from Sigma-Aldrich unless otherwise noted.

Isolation and Culture of Bone Marrow MSCs

Bone marrow MSCs (BM-MSCs) were isolated using a published method [29]. Femurs and tibias were isolated from euthanized ICR mice, and scissors were used to remove epiphyses. The marrow cells were harvested using 23- and 26-gauge hypodermic needles from the diaphyses of femurs and tibias, respectively, followed by flushing with MEM-α supplemented with 5.7 mg/mL NaHCO3, 10% FBS, 100 IU/mL penicillin and 100 mg/mL streptomycin. Cells (2×106 cells/cm2) were added to 60-cm2 culture dishes and incubated as described above. Non-adherent cells were removed by changing the medium every three days. Cells (70% confluent) were removed from the dishes by incubation at 37°C with 0.25% trypsin/EDTA for 5 min and were purified using transient lower-density plastic adherence [29].

Cell Proliferation

To assess the proliferative ability of EGFP-AFSCs, we estimated the population doubling time (PDT) according to the equation: TD = t0 log2/(logNt−logNo), where Nt and No are the numbers of cells harvested and inoculated, respectively, and t is the time of culture (in hours). Cell viability was determined using the trypan blue (#13250, Gibco) exclusion test, and the cells were counted using a hemocytometer [30].

Immunocytochemistry

Cells were fixed with 4% (w/v) PFA for 10 min at room temperature and permeabilized using 0.25% Triton X-100 (w/v) (#0694, Amresco, Solon, OH) for 10 min. To reduce nonspecific antibody-binding, the cells were treated with 0.1% (v/v) Tween-20 (#X251-07, Sigma-Aldrich) in phosphate buffered saline (PBS) (#3070, Amresco) (PBS) containing 3% (w/v) bovine serum albumin (BSA) (#0332, Amresco) for 30 min and then incubated with a rabbit anti-OCT4 antibody (1:200; #SC-9081, Santa Cruz, Dallas, TX) at 4°C overnight. Immune complexes were detected using a goat anti-rabbit IgG secondary antibody labeled with Alex Fluor 394 (#1300; #A11012, Invitrogen, Carlsbad, CA). Nuclei were stained with 0.1 mg/mL 4,6-diamidino-2-phenylindole (DAPI) (#10006; #40043, Biotium, Hayward, CA) for 3 min. The cells were observed using a fluorescence microscope (DMIRB; Leica, Wetzlar, Germany).

Mouse Model of CTx-Induced POF

Six-week-old ICR female mice were administered busulfan (20 mg/kg) (#B2635) and cyclophosphamide (200 mg/kg)
mice, respectively. PBS injection only are designated CTx-AFSCs- and CTx-saline-
mice anesthetized with inhaled isoflurane (#B506, Abbott, Chicago, II). CTx-mice with EGFP-AFSCs transplantation or PBS injection only are designated CTx-AFSCs- and CTx-saline-
mice, respectively.

Cell Transplantation
After six weeks of the POF mouse model established successfully, EGFP-AFSCs were transplanted into the ovaries of CTx-mice using a modification of a published study [31]. In brief, a suspension of 5×10^5 EGFP-AFSCs in 5 μL of sterilized PBS was incubated through a glass pipette (100-μm tip) into each ovary of mice anesthetized with inhaled isoflurane (#M9140, Sigma-Aldrich), and the cells were classified using a standard criteria [34]. An estrus cycle was defined as the period between two successive proestrus phases (proestrus = day 1).

Vaginal Cytology
To validate the reproduction function, the number of estrous cycles will be analyzed by vaginal smear assay every day after one week post AFSCs transplantation and for 14 consecutive days of test. Vaginal cells were collected using a sterile cotton swab, smeared on glass microscope slides, and stained with 0.5% (w/v) methylene blue (#S0389, Sigma-Aldrich), and the cells were classified using a standard criteria [34]. An estrus cycle was defined as the period between two successive proestrus phases (proestrus = day 1).

Mating Trial
Mating trial was initiated after one week post AFSCs transplantation and continued for 24 days. One fertile male was housed with one female mouse from each experimental group. The presence of a copulatory plug was defined as successful mating. Males were randomly rotated among cages after each pregnancy, and the number of offspring per litter was recorded.

Immunohistochemistry
In brief, ovaries were fixed with 4% PFA at 4°C overnight, incubated in 30% sucrose (#S0389, Sigma-Aldrich) at 4°C for three days, frozen, sectioned, incubated with PBS containing 0.25% Triton X-100 for 15 min, and blocked with PBST containing 3% BSA for 30 min. Sections were incubated with a rabbit anti-VASA (DDX4) antibody (1:200; ab13840, Abcam, Cambridge, UK) for 1 h at room temperature, rinsed with PBS, and incubated with a goat anti-rabbit IgG secondary antibody labeled with Alex Fluor 594 (#ab104135, Abcam) on glass slides and observed using a confocal microscope (Leica TCS SP5, Wetzlar, Germany).

Ovarian Morphology and Follicle Counts
The number of follicles in each ovary was estimated according to a modified published protocol [32]. Ovaries were fixed with 4% PFA at 4°C overnight, embedded in paraffin, serially sectioned (8 μm), and stained with hematoxylin-eosin. The number of developing and total healthy follicles were also significantly different. Log-rank (Mantel-Cox) test was used to compare the data for two groups and three groups, respectively. P<0.05 was defined as a statistically significant difference. Log-rank (Mantel-Cox) test was used to compare the percentage of successful mating for three groups and the P<0.01 was defined as a statistically significant difference.

Results
Characterization of EGFP-AFSCs
Most of the EGFP-AFSCs (Figure 1A) strongly expressed EGFP (Figure 1B), and flow cytometric analysis showed that cultured EGFP-AFSCs expressed CD29, CD14, Sca-1, and MHC-I, while did not express MHC-II, hematopoietic lineage markers (CD11b, CD34, CD45) or an endothelial cell marker (CD31) (Figure 1C). Culturing in adipogenic medium induced EGFP-AFSCs to form intracellular lipid droplets (Oil Red O-positive) (Figure 1D). Culturing in osteogenic medium induced EGFP-AFSCs to form calcium deposits and mature nodules structures, which were stained by ARS (Figure 1E). After 21 days of incubation in chondrogenic medium, EGFP-AFSCs stained with toluidine blue exhibited a blue-purple matrix (Figure 1F). These findings showed that AFSCs displayed morphology, immunophenotypes and mesoderm trilineage differentiation capacity similar to MSCs.

To further evaluate the proliferative ability of AFSCs, we performed growth curve and PDT assay compared to BM-MSCs. We found that AFSCs showed higher cell population growth with shorter PDTs than BM-MSCs (31.22±1.10 h and 34.41±0.73 in AFSCs and BM-MSCs, respectively) (Figure 1G). In accordance with previous studies, we confirmed that most of cultured AFSCs expressed OCT4 in the nucleus by immunocytochemistry staining (Figure 1H-J). These findings illustrated that the AFSCs we established exhibited higher proliferation ability than BM-MSCs and expressed OCT4.

Effect of CTx on Ovarian Function and Induced POF
The ovaries of CTx-mice were atrophic compared with those of non-CTx-mice and contained fewer developing follicles at the sixth week post CTx (Figure S1). Further, primordial follicles were significantly fewer in the ovaries of CTx-mice and almost disappeared after six weeks post CTx (Figure 2A). The numbers of developing and total healthy follicles were also significantly

Reverse Transcription-Polymerase Chain Reaction (RT-PCR) Assay
The total RNA was extracted from ovarian tissue using TRIzol reagent (#10296-010, Invitrogen). Reverse transcription reactions were performed using the SuperScript First-Strand Synthesis System (#18064-014, Invitrogen) with 1.6 μg total RNA. PCR was performed with Taq polymerase (#11615-010, Invitrogen). The preceding reactions were performed according to the manufacturers' instructions. PCR reactions were performed as follows: initial denaturation at 94°C for 5 min, 35 cycles of denaturation at 94°C for 30 s, annealing at 55°C for 30 s, elongation at 68°C for 30 s, and final incubation at 72°C for 10 min. The sequences of primers were as follows: forward 5'-AGAGGGAATCTGCGCTGAC-3', reverse 5'-CAAGAAG-GAAGGCTGGGAA-3' for Artb (189 bp); forward 5'-TATAT- CATGGCGCAACAGCA-3'; reverse 5'-GAACTCCAGACAG- GACCAATGT-3' for egfp (219 bp). The expression of β-actin was used as the internal control.

Statistical Analysis
Three biological replicates were performed, and all values represent the mean ± standard error of the mean (S.E.M). An unpaired Student t-test and Tukey's multiple comparison test were used to compare the data for two groups and three groups, respectively. P<0.05 was defined as a statistically significant difference. Log-rank (Mantel-Cox) test was used to compare the percentage of successful mating for three groups and the P<0.01 was defined as a statistically significant difference.

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reduced in the ovaries of CTx-mice, particularly after six weeks (Figure 2B-E). In contrast, the numbers of atretic follicles were not significantly different between the ovaries of CTx- and non-CTx-mice (Figure 2F).

AFSCs Prevent Follicular Atresia and Sustained the Development of Follicles in the Ovaries of CTx-mice

To assess the therapeutic effect of AFSCs, we transplanted EGFP-AFSCs directly into bilateral ovaries of mice with POF six weeks after CTx. Although there were few primordial follicles in the ovaries of CTx-saline- and CTx-AFSCs-mice, during the

Figure 1. Characterization of EGFP-AFSCs in culture. (A) Amniotic fluid stem cells (AFSCs) (third passage) exhibit fibroblast-like morphology and adhere to the surface of plastic cell culture dishes. (B) Harvested AFSCs express enhanced green fluorescence protein (EGFP). (C) Immunophenotypes of AFSCs at their third passage expressed markers of mesenchymal stem cells (MSCs) but not those of hematopoietic and endothelial cells. The black histograms indicate the respective isotype control. (D) Oil Red O staining after 14 days of culture in adipogenic medium showed adipogenic differentiation potential of AFSCs. (E) Alizarin Red S (ARS) staining after 21 days of culture in osteogenic medium showed osteogenic differentiation potential of AFSCs. (F) Toluidine blue O staining after 21 days culture in induction chondrogenic medium showed chondrogenic differentiation potential of AFSCs. Scale bar = 100 µm. (G) Proliferation ability of EGFP-AFSCs (third passage) was better than that of bone marrow MSCs (BMMSCs) (third passage) (n = 3). Data represent the mean ± standard error of mean (S.E.M.) *P<0.05. (H–J) Immunocytochemistry analysis revealed the OCT4 expression (red in I and J) by EGFP-AFSCs with nuclear counterstaining by 4′,6-diamidino-2-phenylindole (DAPI) (blue in H, J and K). Scale bar = 25 µm.

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course of the experiment, the numbers of primordial follicles were significantly higher in CTx-AFSCs-mice during the first four weeks post-transplantation (wpt) (Figure 3A). The results were similar for the numbers of primary and secondary follicles among the three groups (Figure 3B and C). The number of antral follicles of CTx-AFSCs-mice was significantly higher compared with that of CTx-saline-mice, whereas lower compared with that of non-CTx-mice at all times (Figure 3D). The number of total healthy follicles in CTx-AFSCs-mice was higher compared with CTx-saline-mice within 4 wpt and was significantly lower compared with that of non-CTx-mice at all times (Figure 3E). Injection of EGFP-AFSCs significantly decreased the number of atretic follicles up to 4 wpt. In contrast, there was no significant difference between non-CTx- and CTx-saline-mice at all times (Figure 3F). There was no significant difference between the sum of total healthy and atretic follicles of CTx-AFSCs- and CTx-saline-mice (Figure 3G). These results indicated that AFSCs could transiently prevent follicular atresia and sustain the follicular development in the ovaries of mice with POF.

AFSCs Restore the Fertility of Mice with POF

The number of estrous cycles of CTx-AFSCs-mice (3.4±0.22) was significantly higher compared with that of CTx-mice (0.4±0.22) but not significantly different compared with that of non-CTx-mice (4±0.28) (Figure 4A). Although the CTx-AFSCs-mice reached 100% successful mating rate significantly slower than non-CTx control group (P<0.01), the CTx-saline-mice reached only 40% successful mating in the 24-day test period and was significantly worse than CTx-AFSCs-mice (P<0.01) (Figure 4B). Moreover, consistent with the numbers of antral follicles, the average number of litters produced by CTx-AFSCs-mice (6.9±0.85) was significantly higher compared with those of CTx-mice (0) and lower compared with those of non-CTx-mice (13.2±1.09) (Figure 4C).
Figure 3. Quantitative analysis of ovarian follicles of CTx-mice transplanted with EGFP-AFSCs. (A–F) (A) Primordial, (B) primary, (C) secondary, (D) antral, (E) total healthy, and (F) atretic follicles were counted in the mice ovaries. (G) The transplantation of EGFP-AFSCs did not alter the number of the sum of total healthy and atretic follicles in the ovaries of CTx-mice. Con = Non-CTx-mice. Saline = CTx-saline-mice. AFSCs =
EGFP-AFSCs do not Contribute to the Germline in Ovaries of CTx-Mice

The progeny of CTx-AFSCs-mice appeared normal (Figure 4D and E), and none expressed EGFP (Figure 4F), indicating that offspring were derived from the recipient’s germline (n = 69). Therefore, we tracked the fate of the transplanted EGFP-AFSCs within the ovaries of recipients. EGFP-AFSCs were detected in ovarian tissue at 1 wpt and persisted for at least four weeks (Figure 5A-D). Neither the germ cell marker VASA [35] (Figure 5D) nor the granulosa cell marker follicle stimulating hormone receptor (FSHR) [36] (data not shown) was detectably expressed by EGFP-AFSCs. Moreover, the level of the mRNA encoding EGFP in the ovaries of CTx-AFSCs-mice decreased gradually after transplantation, and little was detected after 5 wpt (Figure 5E).

Discussion

Cancer CTx using busulfan/cyclophosphamide are designed to destroy highly proliferating cells by inducing cross-linking of DNA [37], which affect mitochondria and result in the activation of apoptotic pathway [38]. Accordingly, it is widely believed that chemotherapeutic drugs may massively eliminate granulosa cells [39], which are required for oocyte survival and follicle development [40]. The ensuing damage to growing follicles by CTx increases recruitment of primordial follicles and, in turn, the number of all types of follicles reduced [41]. Consistent with this hypothesis and previous studies [42,43], our results showed that CTx treatment significantly reduced various types of follicles and number of total healthy follicles (Figure 2A-E). Nonetheless, although CTx largely diminished number of total healthy follicles, the number of atretic follicles was not significantly different at any point during the 6 weeks observation (Figure 2F). According to the previous studies, chemotherapeutic drugs significantly increased follicle atresia predominantly within 72 h after the treatment and the number of atretic follicle quickly returned to the basal level within one week after the treatment. Subsequently, the degenerative follicles were removed from the ovary within a 3-day period [32,44]. This should be the reason of the discordance in the observed numbers of follicle loss versus atresia. Consequently, the exhaustion of primordial follicles and continued follicular atresia severely impaired the ovarian function and resulted in POF after six weeks of CTx.

In this study, we demonstrated that AFSCs increased healthy ovarian follicles and restored fertility in a mouse model system of POF induced by CTx. In principle, AFSCs may sustain the number of healthy ovarian follicles in POF mice either by promoting de novo folliculogenesis or by inhibiting follicular atresia.

Figure 4. Reproductive function was improved in CTx-mice transplanted with EGFP-AFSCs. (A) The number of estrous cycles in two weeks was evaluated in the mice. Con = Non-CTx-mice (n = 5). Saline = CTx-saline-mice. (n = 5). AFSCs = CTx-AFSCs-mice. (n = 5). Data represent the mean ± S.E.M. Lower case letters (a, b, and c) represent significant differences (P<0.05) among groups. (B) The successful mating rate was assessed in a 24-day period. The presence of a copulatory plug indicated successful mating. Con = Non-CTx-mice. (n = 10). Saline = CTx-saline-mice. (n = 10). AFSCs = CTx-AFSCs-mice. (n = 10). Log-rank (Mantel-Cox) test was used to compare the data among groups. P<0.01 represents significant differences between groups. (C) The litter numbers were counted. Con = Non-CTx-mice (n = 10). Saline = CTx-saline-mice. (n = 4). AFSCs = CTx-AFSCs-mice (n = 10). Data represent the mean ± S.E.M. Lower case letters (a, b, and c) represent significant differences (P<0.05) among groups. (D–F) The F1 litters showed normal gross appearance with no progeny bearing EGFP (arrow) as compared to the EGFP transgenic mice (arrowhead). doi:10.1371/journal.pone.0106538.g004
AFSCs Prevent Follicle Atresia and Recover the Fertility

AFSCs served as an internal control. W = Weeks. (Neg) was used as a positive and negative control respectively. Tissues harvested from an EGFP mouse (Pos) and a CTx-saline-mouse ovaries at different times after EGFP-AFSCs transplantation. Ovarian transcription polymerase chain reaction (RT-PCR) analysis of recipient at 4 weeks post-transplantation. Scale bar = 50 μm.

We further showed a significant decrease in the number of atretic follicles in the ovaries of CTx-AFSCs-mice that persisted for three (C) weeks of transplantation. DAPI (blue) was used to stain nuclei. Scale bar = 50 μm. (D) Analysis of VASA (red) and EGFP (green) marker in vitro [47], and therefore it might be possible that the transplanted AFSCs expressed oocyte marker in vitro [47], and therefore it might be possible that the transplanted AFSCs have the potential to differentiate into primitive oocyte and participate in the following folliculogenesis within ovaries of POF mice. However, we found no progeny of CTx-AFSCs-mice derived from the EGFP-AFSCs and none expressed EGFP. Moreover, transplanted EGFP-AFSCs were distributed within the ovarian interstitium rather than follicles and expressed neither a marker of differentiated germ cells (Figure 5D) distributed within the ovarian interstitium rather than follicles and expressed neither a marker of differentiated germ cells (Figure 5D) nor of granulosa cells (data not shown). These results indicated that the transplanted AFSCs per se did not participate in the folliculogenesis.

We further showed a significant decrease in the number of atretic follicles in the ovaries of CTx-AFSCs-mice that persisted for four weeks (Figure 3F), which is in correlation with the existence period of EGFP-AFSCs (Figure 5E). Notably, the numbers of total healthy and atretic follicles were not significantly different between CTx-AFSCs-mice and mice that were not engrafted (Figure 3G). These data strongly suggested that AFSCs exerted therapeutic effect on increasing the number of healthy ovarian follicles by preventing follicular atresia.

Supporting Information

Figure S1 Appearance and histological analyses of ovaries of CTx-mice transplanted with EGFP-AFSCs. (A–C) Transplanted EGFP-AFSCs (green) can be observed in the ovaries after one (A), two (B), and three (C) weeks of transplantation. DAPI (blue) was used to stain nuclei. Scale bar = 50 μm. (D) Analysis of VASA (red) and EGFP (green) expression showed that there is no co-localization in recipient ovaries at 4 weeks post-transplantation. Scale bar = 50 μm. (E) Reverse transcription polymerase chain reaction (RT-PCR) analysis of recipient ovaries at different times after EGFP-AFSCs transplantation. Ovarian tissues harvested from an EGFP mouse (Pos) and a CTx-saline-mouse (Neg) was used as a positive and negative control respectively. β-actin mRNA served as an internal control. W = Weeks. doi:10.1371/journal.pone.0106538.g005

Collectively, our findings demonstrated that AFSCs may prevent follicle atresia in the ovaries of mice with POF and restore fertility. These findings suggest the potential of AFSCs to treat infertility.

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Author Contributions
Conceived and designed the experiments: GYX IHL C. Cheng C. Chang
WTKC SCW. Performed the experiments: GYX SCW. Analyzed the data: GYX C. Chang YHL. Contributed reagents/materials/analysis tools: GYX SCW. Contributed to the writing of the manuscript: GYX IHL C. Cheng.

References
1. Jemal A, Siegel R, Ward E, Murray T, Xu J, et al. (2007) Cancer statistics, 2007. CA Cancer J Clin 57: 43–66.
2. Chemaitilly W, Mertens AC, Mitby P, Whitton J, Stovall M, et al. (2006) Acute ovarian failure in the childhood cancer survivor study. J Clin Endocrinol Metab 91: 1723–1728.
3. Byrne J (1999) Long-term genetic and reproductive effects of ionizing radiation and chemotherapeutic agents on cancer patients and their offspring. Teratology 59: 210–219.
4. Hudson MM (2010) Reproductive outcomes for survivors of childhood cancer. Obstet Gynecol 116: 1117–1183.
5. Oka Y, Ohba E, Kuroki T, Kojima S, Kiyonari S, et al. (2007) A novel mouse model of chemotherapeutic-induced ovarian failure. J Clin Endocrinol Metab 25: 100–106.
6. Bruning PF, Pit MJ, de Jong-Bakker M, van den Ende A, Hart A, et al. (1990) Bone marrow density after adjuvant chemotherapy for premenopausal breast cancer. Br J Cancer 61: 308–310.
7. Carter J, Rowland K, Chi D, Brown C, Abu-Rustum N, et al. (2005) Gynecologic cancer treatment and the impact of cancer-related infertility. Gynecol Oncol 97: 90–95.
8. Lee BJ, Selemeni K, Nikura Y, Nikura T, Klein R, et al. (2007) Bone marrow transplantation generates immature oocytes and rescues long-term fertility in a preclinical mouse model of chemotherapeutic-induced premature ovarian failure. J Clin Endocrinol Metab 25: 3198–3204.
9. Takehara Y, Yabuuchi A, Ezoe K, Kuroda T, Yamadera R, et al. (2013) The restoration of effects of adipose-derived mesenchymal stem cells on damaged ovarian function. Lab Invest 93: 181–193.
10. Fu X, He Y, Xie C, Liu W (2008) Bone marrow mesenchymal stem cell transplantation improves ovarian function and structure in rats with chemotherapeutic-induced ovarian damage. Cytotherapy 10: 355–363.
11. Basset MA, Wynn RF, Jowit SN, Waddell JI, Fairbairn LJ, et al. (2004) Study of telomere length reveals rapid aging of human marrow stromal cells following in vitro expansion. Stem Cells 22: 675–682.
12. Bryant-Greenwood GD (1998) The extracellular matrix of the human menstrual cycle: structure and function. Placenta 19: 1–11.
13. De Coppi P, Bartsch G, Jr., Siddiqui MM, Xu T, Santos CC, et al. (2007) Bone marrow adipose tissue mesenchymal stem cells are capable of converting to smooth muscle cells in the cryo-injured rat bladder and prevent compensatory hypertrophy of surviving smooth muscle cells. J Urol 177: 369–376.
14. De Gemmis P, Lorandi C, Gentili C, van Griensven M, et al. (2012) Isolation of mesenchymal stem cell lines with potential for therapy. Nat Biotechnol 25: 100–106.
15. Niwa H, Yamamura K, Miyazaki J (1991) Efficient Selection for High-Expression Transfectants with a Novel Eukaryotic Vector. Gene 108: 193–199.
16. Peister A, Mellad JA, Larsson BL, Hall BM, Gibson LF, et al. (2004) Adipose stem cells from bone marrow (MSCs) isolated from different strains of inbred mice vary in surface epitopes, rates of proliferation, and differentiation potential. Blood 103: 1662–1668.
17. Johnstone B, Hering TM, Caplan AI, Goldberg VM, Yao JU (1998) In vitro chondrogenesis of bone marrow-derived mesenchymal progenitor cells. Exp Cell Res 238: 265–272.
18. BC Cheng, Lian WS, Hsiao FS, Liu IH, Lin SP, et al. (2012) Isolation and characterization of novel murine epiphysis derived mesenchymal stem cells. PLoS One 7: e36085.
19. S. Naderi, S. Soleimani, M. Hosseini RH, Massumi M, Atashian A, et al. (2007) An efficient method for isolation of murine bone marrow mesenchymal stem cells. Int J Dev Biol 51: 723–729.
20. Zou K, Yuan Z, Yang Z, Luo H, Sun K, et al. (2009) Production of offspring from a germine stem cell line derived from neonatal ovaries. Nat Cell Biol 11: 631–636.
21. Johnson J, Canning J, Kaneko T, Prö JK, Tilly JL (2004) Germine stem cells and follicular renewal in the postnatal mammalian ovary. Nature 428: 145–150.
22. Morita Y, Perez GI, Mazarei DV, Tilly KJ, Tilly JL (1999) Targeted expression of Bcl-2 in mouse oocytes inhibits ovarian follicle atresia and prevents spontaneous and chemotherapeutic-induced oocyte apoptosis in vivo. Mol Endocrinol 13: 841–850.
23. Nelson JF, Felician LS, Randall PK, Sims C, Finch CE (1982) A longitudinal study of castrate cyclicity in aging C57Bl/JJ mice I. Cycle frequency, length and vaginal cytology. Biol Reprod 27: 327–339.
24. Tomooka Y, Tsunamiwa N, Takahashi Y, Matsui Y, Sato M, et al. (2000) Expression and intracellular localization of mouse Vasa-homologue protein during germ cell development. Mech Dev 93: 139–149.
25. Hillier SG (2001) Gonadotropic control of ovarian follicular growth and development. Mol Cell Endocrinol 179: 39–46.
26. Helløedt T, Petermann E, Lundin C, Hodgson B, Sharma RA (2000) DNA repair pathways as targets for cancer therapy. Nature Reviews Cancer 8: 193–204.
27. Zhao XJ, Huang YH, Yu YC, Xin XY (2010) GoRH antagonist cetrorelix inhibits mitochondria-dependent apoptosis triggered by chemotherapy in granulosa cells of rats. Fertil Steril 93: 69–75.
28. Desmeules P, Devine JJ (2006) Characterizing the ovootoxicity of cyclophosphamide metabolites on cultured mouse ovaries. Toxicol Sci 90: 500–509.
29. Matsuk MM, Burns KH, Viveiros MM, Eppig JJ (2002) Intracellular communication in the mammalian ovary: oocytes carry the conversation. Science 296: 2170–2180.
30. Rosendahl M, Andersson CY, la Cour Freiesleben N, Juul A, Lossl K, et al. (2010) Dynamics and mechanisms of chemotherapeutic-induced ovarian follicle depletion in women of fertile age. Fertil Steril 94: 156–166.
31. Beck-Peccoz P, Persani L (2006) Premature ovarian failure. Orphanet J Rare Dis 1: 9.
32. Groswain D, Conway GS (2005) Premature ovarian failure. Hum Reprod Update 11: 391–410.
33. Perrillo SK, Desmeules P, Truong TQ, Devine JJ (2011) Detection of DNA damage in oocytes of small ovarian follicles following phosphoramidate mustard exposures of cultured rodent ovaries in vitro. Toxicology and Applied Pharmacology 253: 94–102.
34. McLaren A (2003) Primordial germ cells in the mouse. Dev Biol 262: 1–15.
35. Pepling ME (2006) From primordial germ cell to primordial follicle: mammalian female germ cell development. Genesis 44: 622–632.
36. Lai D, Wang F, Chen Y, Wang L, Yang Y, et al. (2013) Human amniotic fluid stem cells have a potential to recover ovarian function in mice with chemotherapy-induced sterility. BMC Dev Biol 13: 34.
37. Hughes FM Jr, Gorospe WC (1991) Biochemical identification of apoptosis (programmed cell death) in granulosa cells: evidence for a potential mechanism underlying follicular atresia. Endocrinology 129: 2413–2422.
38. Kaipa A, Huch AJ (1997) Regulation of ovarian follicle atresia. Ann Rev Physiol 59: 349–363.
39. Jiang JY, Cheung CK, Wang Y, Tsang BK (2003) Regulation of cell death and cell survival gene expression during ovarian follicular development and atresia. Front Biosci 8: 2222–237.
40. Yoon BS, Moon JH, Jun EK, Kim J, Maeng I, et al. (2010) Secretory profiles and wound healing effects of human amniotic fluid-derived mesenchymal stem cells. Stem Cells Dev 19: 897–902.
41. Miranda-T, Hartinger J, Loroleti C, Gentili C, van Griensven M, et al. (2012) Prionogenic soluble factors from amniotic fluid stem cells mediate the recruitment of endothelial progenitors in a model of ischemic fasciocutaneous flap. Stem Cells Dev 21: 2179–2186.
53. Cheng FC, Tai MH, Sheu ML, Chen CJ, Yang DY, et al. (2010) Enhancement of regeneration with glia cell line-derived neurotrophic factor-transduced human amniotic fluid mesenchymal stem cells after sciatic nerve crush injury. J Neurosurg 112: 868–879.

54. Knight PG, Glister C (2006) TGF-beta superfamily members and ovarian follicle development. Reproduction 132: 191–206.

55. Knight PG, Glister C (2003) Local roles of TGF-beta superfamily members in the control of ovarian follicle development. Anim Reprod Sci 78: 165–183.

56. Shin SY, Lee JY, Lee E, Choi J, Yoon BK, et al. (2006) Protective effect of vascular endothelial growth factor (VEGF) in frozen-thawed granulosa cells is mediated by inhibition of apoptosis. Eur J Obstet Gynecol Reprod Biol 125: 233–238.

57. Linher-Melville K, Li J (2013) The roles of glial cell line-derived neurotrophic factor, brain-derived neurotrophic factor and nerve growth factor during the final stage of folliculogenesis: a focus on oocyte maturation. Reproduction 145: R43–54.