Mesenchymal stem cells reportedly have a marked effect on tumor growth or suppression. However, it remains uncertain whether adipose-derived mesenchymal stem cells (ADSCs) from grafted fat can contribute to breast cancer growth and recurrence. In the present study, interactions between ADSCs and MCF-7 breast cancer cells were evaluated in a Matrigel co-culture system and in an in vivo nude mouse model. Results suggested that MCF-7 cells exerted tumor tropism effects on ADSCs and this may be regulated by chemokines, such as the macrophage inflammatory protein (MIP)-1γ and MIP-3α. Additionally, ADSCs significantly induced tumorsphere formation in vitro and promoted tumorigenicity in vivo. RT-qPCR analysis indicated that tumorsphere formation by MCF-7 cells was associated with the induction of stem-like properties, which was mediated by epithelial-mesenchymal transition. Together, the present findings indicated that ADSCs exhibit tropism and induce tumorsphere formation of MCF-7 cells.

Introduction

Breast cancer remains one of the most frequent malignancies in women, accounting for ~350,000 annual mortalities in recent years worldwide (1,2). Surgical treatment is currently the preferred option for almost all types of breast cancer. However, surgical incision procedures lead to the loss of breast volume and distortion of shape, and follow-up radiation therapy often results in breast tissue fibrosis and poor wound healing (3,4). Among the plastic surgery techniques currently available to reconstruct the breast, fat grafting is gaining major interest given its ease to harvest, low morbidity and capacity to improve the tissue quality, particularly in breast-conserving surgery (5-9). However, the possibility that breast cancer cells may still reside in patients with breast cancer after the surgical treatment cannot be completely excluded.

Adipose tissue is a multifunctional organ mainly consisted of mature adipocytes and the adipose mesenchymal stromal cells (MSCs)/stromal vascular fraction. Adipose MSCs are heterogeneous and contain several populations, including adipose-derived stem cells (ADSCs), endothelial progenitor cells, pre-adipocytes, lymphocytes, mast cells, pericytes, and adipose-resident macrophages (10,11). Transplantation of adipose tissue, consisting of ADSCs that are metabolically active and secrete various cytokines, does not simply behave as an inert filler but it tends to influence the cancer microenvironment (12,13). Hence, whether fat grafting can be applied to patients following breast cancer surgery is still a controversial issue.

It is widely recognized that multipotent ADSCs with their regenerative features, such as pro-angiogenic, anti-apoptotic, pro-proliferative and multipotent differentiation characteristics, within the transferred fat mainly contribute to the restorative and reconstructive qualities of autologous fat grafting (14-16). Unfortunately, these regenerative features are also assumed to be associated with tumor initiation and metastasis, causing safety concerns in clinical utilization. Notably, the majority of studies investigate the interaction between ADSCs and breast cancer cells and are performed in a two-dimensional (2D) context in vitro (17-20), which does not fully recapitulate the in vivo condition. Hence, an in vitro 3D culture system was established and applied an in vivo animal model to investigate the interaction between ADSCs and MCF-7 breast cancer cells in tumor development in the present study, mainly focusing on the tropism of ADSCs towards the breast cancer cells and the potential mechanism of ADSCs on promoting MCF-7 cells progression.

Materials and methods

Correspondence to: Dr Jianhua Gao or Dr Feng Lu, Department of Plastic and Cosmetic Surgery, Nanfang Hospital, Southern Medical University, 1838 Guangzhou North Road, Guangzhou, Guangdong 510515, P.R. China
E-mail: gaopsn@163.com
E-mail: doctorlufeng@hotmail.com

*Contributed equally

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Medical University Institutional Review Board (Guangzhou, China) and the patient provided written informed consent to donate remaining tissues after liposuction. All procedures performed involving animal experiments were approved by the Nanfang Hospital animal ethic committee (permit no. NFYY201679) and was conducted in accordance with the ethical standards of the National Health and Medical Research Council China.

**Cell preparation and identification.** Human ADSCs were isolated from abdominal liposuction aspirates of a 28-year-old female patient during an abdominoplasty procedure with informed consent under approval from the Southern Medical University Institutional Review Board. Briefly, fat aspirate was washed with PBS, centrifuged at 800 x g at 25°C for 5 min and digested with 0.1% collagenase at 37°C for 2 h. The dispersed material was centrifuged (170 x g; 25°C) for 5 min, and the pellet was resuspended in Dulbecco's modified Eagle's medium (DMEM; Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA) with 10% fetal bovine serum (FBS; Gibco; Thermo Fisher Scientific, Inc.), 100 U/ml penicillin and 100 µg/ml streptomycin, and seeded in flasks. Next day, non-adherent cells were removed, and the remaining cells were cultured until 80% confluency. Passage 3 ADSCs were used in the following experiments. For the senescence evaluation of used cells, passage 3 ADSCs were further subjected to replicative senescence experiments. For a control culture, the same senescence experiments were conducted on ADSCs at passage 10.

MCF-7 cells were obtained from the Research Laboratory Collaboration Alliance of Nan Fang Hospital (Guangzhou, China). All cells used in the present study were maintained in DMEM supplemented with 10% FBS, 100 U/ml penicillin and 100 µg/ml streptomycin, and seeded in flasks. Next day, non-adherent cells were removed, and the remaining cells were cultured until 80% confluency. Passage 3 ADSCs were used in the following experiments. For the senescence evaluation of used cells, passage 3 ADSCs were further subjected to replicative senescence experiments. For a control culture, the same senescence experiments were conducted on ADSCs at passage 10.

To induce multilineage differentiation, ADSCs were cultured in adipogenic, osteogenic, and chondrogenic medium as previously described (21). Fat, bone and cartilage cells differentiated from ADSCs were identified by staining with Oil Red O (15 min at 25°C), Alizarin red (5 min at 25°C) or Alcian blue (30 min at 25°C), respectively.

**Senescence-associated β-galactosidase assay.** β-Galactosidase assay was used for assessing senescence of used cells using a Senescence-associated β-galactosidase Staining kit (cat. no. C0602; Beyotime Institute of Biotechnology, Haimen, China) as previously described (2,23). Briefly, passage 3 and 10 ADSCs were washed in PBS, fixed for 10 min (room temperature) in 2% formaldehyde, washed, and incubated with the working solution containing 0.05 mg/ml 5-bromo-4-chloro-3-indoly-l-b-d-galactopyranoside (X-gal). After incubation at 37°C for 12 h in the dark, the nucleus was counterstained with nuclear fast red (cat. no. N8002; Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) and positive cells were observed under a light microscope at x200 magnification. The percentage of senescent cells was calculated by the number of blue, β-galactosidase-positive cells out of all cells in 6 different microscope fields. Senescence assays were performed in triplicate.

**Preparation of co-culture conditioned media.** To study the effects of cytokines from a co-culture system on MCF-7 cells, ADSCs and MCF-7 co-culture conditioned media (AM-CM) was prepared. The same amount (4x10^6) of ADSCs and MCF-7 cells were plated in a flask and co-cultured to 80% confluency. Serum-free DMEM was added to the flask and cultured for 48 h at 37°C after being washed with PBS twice. The AM-CM was filtered and stored at -80°C for a week, until further use.

**Cell membrane labeling and co-culture in Matrigel.** To track the interaction between cells, ADSCs and MCF-7 cells were stained with Vybrant<sup>®</sup> Di Cell-Labeling Solution and DiO Cell-Labeling Solution, respectively (Invitrogen; Thermo Fisher Scientific, Inc.), according to the manufacturer's protocol. The same amount (4x10^6) of ADSCs and MCF-7 cells were mixed uniformly and seeded in Growth-factor-reduced Matrigel (cat. no. 356230; BD Biosciences, Franklin Lakes, NJ, USA) to fabricate a 3D culture system. The interactions between ADSCs and MCF-7 cells was observed continuously in Matrigel for 96 h at 37°C and 5% CO<sub>2</sub>; using a confocal laser-scanning microscope (FV10i-W; Olympus Corporation, Tokyo, Japan). Co-culture assays were performed in quadruplicate.

**Scanning electron microscopy (SEM).** For scanning electron microscopy, the same amount (4x10^6) of ADSCs and MCF-7 cells were co-cultured at 37°C in Matrigel on round glass coverslips (Carl Roth GmbH & Co. KG, Karlsruhe, Germany) in 12-well plates. After 2 days, co-culture samples were fixed in PBS (1 ml) with 2% glutaraldehyde and incubated at 25°C for 60 min. Samples were dehydrated in increasing concentrations of acetone, critical-point dried, fixed to stubs with colloidal silver, sputtered with gold using a MED 010 coater and examined under a S-3000N scanning electron microscope (Hitachi Ltd., Tokyo, Japan). An acceleration voltage of 20 kV was used, and images were observed using S-3000N scanning electron microscope (Hitachi, Ltd.).

**In vitro Transwell migration assay.** Adipose stromal cells (ASC) migration assays were performed in triplicate using Transwell migration chambers (8-µm pore size; BD Biosciences). ADSCs (3x10^5) were plated in the upper wells, whereas MCF-7 cells (3x10^6) or AM-CM (600 µl) were dispensed in the lower chamber. Controls were represented by serum-free DMEM. After 24 h of incubation at 37°C, cells that remained on the top of the filter were scrubbed off, and cells that had migrated to the underside of the filter were fixed in methanol and stained with DAPI for 30 min at 25°C. Migrated cells were manually counted under a fluorescence microscope. Migration assays were performed in triplicate.

To further investigate the mechanisms underlying the cells migration in this co-culture system, MCF-7 cells (1x10^6) were co-cultured with ADSCs cells (1x10^6) in Matrigel. MCF-7 alone served as the control. Cells in both groups were collected for gene analysis at day 1, 5, 9.

**Tumorsphere formation in vitro.** To investigate the tumoursphere formation capacity of MCF-7 cells under the influence of either contact or secretion signals of ADSCs, MCF-7 cells (1x10^6) were either co-cultured with ADSCs (1x10^6) or treated with 1 ml AM-CM in Matrigel in 6-wells plates. MCF-7 cells (1x10^6) alone served as the control. The diameters of tumourspheres in four random fields of each well were counted and
imaged. Tumorsphere formation assays were performed in triplicate.

To further investigate the effects of cytokines from a co-culture system on tumorsphere formation of MCF-7 cells, MCF-7 cells (1x10⁵) were treated with 1 ml AM-CM (replaced every 3 days) in Matrigel in 6-wells plates. MCF-7 cells (1x10⁵) alone served as the control. On day 1, 5, and 9, the Matrigel cultures were made into single-cell suspensions using Dispase (10 mg/ml; Roche Diagnostics, Basel, Switzerland) and cells in both groups were collected for subsequent gene analysis.

Reverse transcription-quantitative polymerase chain reaction (RT-qPCR). The co-cultures and MCF-7 cells alone that were cultured on Matrigel were made into single-cell suspensions using Dispase (10 mg/ml; Roche Diagnostics). Sample RNA was extracted using TRIzol® Reagent (Invitrogen; Thermo Fisher Scientific, Inc.) and used for cDNA synthesis with the DBI-2220 Bestar® qPCR Primer Array (DBI Bioscience, Rockville, MD, USA) and the ABI 7500 Fast Real-Time PCR system (Applied Biosystems; Thermo Fisher Scientific, Inc.). The primers used in the present study were: Macrophage inflammatory protein (MIP)-1β, forward 5'-CCACTCACACATACTGCTCTTA-3', and reverse 5'-AGTGTAGGAACCCTGCAATTAC-3'; MIP-3α, forward 5'-CCAAAGACTGGGTACTCAACA-3', and reverse 5'-GAGTAGACACTGACTGATCACCATTAA-3'; SOX2, forward 5' -GAG AGA GAA AGA AAG CAT-3'; MIP-3α, forward 5'-GAGAGACACTGACTGATCACCATTAA-3'; Vimentin, forward 5'-GAT TCA CTC CCT CTG GTT GAA TCA-3'; OCT4, forward 5'-GAGAGACACTGACTGATCACCATTAA-3'; and reverse 5'-CTCTCCTCAGGTTCCTGCTA TACT-3'; E-Cadherin, forward 5'-CTCGACACCACCGATTAATG-3', and reverse 5'-CCAGGCCTAGACCAAGAAAT-3'; Vimentin, forward 5'-GATTCCTCCCTTCTGTTGATAC-3', and reverse 5'-GTACATCGTGATCTGAGAATG-3'; and β-actin, forward 5'-GGACCTGACTGACTCTCAT-3', and reverse 5'-CGTACGCCAGCTTCTCCTTAAT-3'. Thermocycling conditions were as follows: 95°C for 10 min; followed by 40 cycles of 10 sec at 95°C and 20 sec at 55°C. PCR specificity was assessed by the 2²ΔΔCq method (24); β-actin was used as an endogenous reference gene and for normalization.

Western blot analysis. The Matrigel cultures were made into single-cell suspensions using Dispase (10 mg/ml; Roche Diagnostics). Protein was extracted from cells (1x10⁵) using M-PER Mammalian Protein Extraction Reagent (Thermo Fisher Scientific, Inc.) and quantification of protein lysates was conducted with the Bradford method. Protein products (60 µg/lane) were separated using 10% SDS-PAGE and subsequently transferred overnight onto a polyvinylidene difluoride membranes (EMD Millipore, Billerica, MA, USA). Membranes were blocked in 5% milk for 1 h at 25°C and incubated with the following primary antibodies: Anti-E-Cadherin [1:500; Cell Signaling Technology (CST), Inc., Danvers, MA, USA; cat. no. 3195]; or anti-Vimentin antibody (1:500; CST, Inc.; cat. no. 5741). The membranes were cultured with horseradish peroxidase-conjugated anti-rabbit immunoglobulin G secondary antibody (1:10,000; cat. no. 111-035-003; Jackson ImmunoResearchLaboratories, Inc., West Grove, PA, USA) for 30 min at 25°C, and proteins were visualized with the WesternBreeze Chemiluminescent Detection kit (Thermo Fisher Scientific, Inc.). β-actin (1:1,000; cat. no. 4970; CST, Inc.) served as an internal control.

Tumor-bearing mice preparation and in vivo imaging of ADSCs homing to tumors. Female nude mice (n=9; age, 4-5 weeks; weight, 13-15 g) were purchased from the Southern Medical University Laboratory Animal Center and were maintained in microisolator cages at the Animal Experiment Center of Nanfang Hospital. Mice were housed in a pathogen-free animal facility (25±2°C; 55% humidity) with ad libitum access to standard food and water, and a 12-h light/dark cycle. Tumor-bearing models were prepared as previously described (25). Briefly, MCF-7 cells (2x10⁵), resuspended in 0.5 ml PBS, were subcutaneously injected into each side of groin adipose pad of the nude mouse. Tumor formation was observed at the groin area 2-3 weeks following the MCF-7 cells injection and the tumor-bearing mice were used in subsequent in vivo experiments. A cell tracer Vybrant® DiI was used to trace ADSCs in vivo; DiI labeling solution was prepared following the manufacturer's protocol. Briefly, 5x10⁵ ADSCs in 1 ml of PBS were mixed with 5 µl of DiI labeling solution and incubated for 15 min at 37°C. After labeling, cells were washed and injected into the tail veins of tumor-bearing mice. Bioluminescence imaging of mice and excised organs (lung, heart, liver and kidney) and tumors was conducted to trace the ADSCs using an in vivo Multispectral Imaging Systems FX, bioluminescence imaging system (Carestream Health, Inc., Rochester, NY, USA) at 4 weeks following ADSC injection.

Xenograft assays in nude mice and frozen sectioning. Female nude mice (age, 4-5 weeks; weight, 13-15 g; n=30) were purchased from the Southern Medical University Laboratory Animal Center and were maintained as aforementioned. For xenograft experiments, MCF-7 cells (1x10⁶) mixed with ADSCs (1x10⁶) were injected into the right groin adipose pad, whereas MCF-7 cells (1x10⁵) alone injected into the left groin adipose pad served as a control. Tumor volumes were measured using Vernier calipers every 4 days for 32 consecutive days and calculated using the following formula: Tumor volume = (length x width²)/2, following previously published protocols (26,27). The mice were sacrificed at 32 days post-inoculation by cervical dislocation after being anesthetized by 1% pentobarbital (40 mg/kg), and the tumor samples were removed for further analysis.

For frozen sections, fresh excised tumor samples were embedded in frozen section embedding medium optimal cutting temperature (OCT) compound and were frozen rapidly at -20°C. The frozen samples were then subjected to hematoxylin and eosin (H&E) staining, following a previously published protocol (28). Briefly, the frozen samples were cryosectioned into 5-µm slices and mounted onto charged microscope slides. The slices were washed with distilled water and 30% isopropanol to remove the OCT compound, and stained with the H&E working solution (hematoxylin for 4 min and eosin for 30 sec) at 25°C. Following staining, the sections were mounted with neutral balata (Nanjing KeyGen Biotech Co., Ltd., Nanjing, China) for 30 min at 25°C and observed under a light microscope.
Statistical analysis. Quantitative results were expressed as the means ± standard deviation. The comparison between co-culture or AM-CM treated samples at different time points with MCF-7 cells was examined using the Student’s t-test with SPSS 22.0 software (IBM Corp., Armonk, NY, USA). Multiple comparisons of migrated ADSCs in Transwell migration assay and tumorspheres diameter in the tumorsphere formation assays were performed using one-way analysis of variance and the least significant difference post hoc analysis was performed to ascertain significance between groups. P<0.05 was considered to indicate a statistically significant difference.

Results

ADSCs characterization and senescence evaluation. The isolated ADSC cells expressed lipid droplets, matrix mineralization, cartilage-specific proteoglycans and were positive for Oil Red O (Fig. 1A), Alizarin red (Fig. 1B), and Alcian blue (Fig. 1C). Senescence degree of ADSCs was evaluated by in situ senescence-associated-β-galactosidase assay. Few senescent cells were observed in passage 3 ADSCs (Fig. 1D), whereas numerous senescent cells were observed in passage 10 ADSCs (Fig. 1E). Quantification analysis revealed that there was a significantly decreased percentage of blue, β-galactosidase-positive cells in passage 3 compared with that in the passage 10 ADSCs (Fig. 1F).

Tumor tropism of ADSCs, chemokine expression and cell interactions. ADSCs and MCF-7 cells were evenly distributed at 12 h after co-culture (Fig. 2A). At 24 h, ADSCs were touching and surrounding MCF-7 cells (Fig. 2B). At 96 h, ADSCs were observed inside MCF-7 tumorspheres. Experiments were repeated 4 times. Scale bar, 100 µm. ADSCs, adipose-derived mesenchymal stem cells.
bioluminescence imaging live fluorescence system and subsequently the tumors and important organs (lung, heart, liver and kidney) were carefully dissected from mice and observed under the same system. The results revealed that the fluorescence signal was mainly concentrated in tumor tissue and nearly no signal was observed in other important organs (Fig. 4A and B). H&E staining and Dil-labelled ADSCs were observed in sections (Fig. 4C and D, respectively). Collectively, these
results suggested that ADSCs mainly migrated to tumor site following intravenous injection, possibly through the circulatory system.

The morphological appearance of co-cultures of ADSCs and MCF-7 cells was observed by a scanning electron microscope. Spherical MCF-7 cells grew into tumorspheres with irregular surfaces in Matrigel substrate (Fig. 5A). Tumorspheres were surrounded by ADSCs, which were recognized by their fltly spread morphology (Fig. 5B). At higher magnification, the construction of connections between ADSCs and MCF-7 cells were observed (Fig. 5C).

ADSCs enhance tumorsphere formation, cancer stem cell (CSC) marker expression, and in vivo tumor formation. Tumorsphere formation, which is a property of cancer stem cells, was observed in our 3D culture system. MCF-7 cells co-cultured with ADSCs exhibited a significant capacity to form tumorspheres, whereas the AM-CM also exhibited a marked effect on tumorsphere formation (Fig. 6A). Quantitative analysis revealed that tumorspheres in co-culture groups (ADSCs + MCF-7) were larger than those treated with AM-CM (MCF-7/AM-CM). As a control, MCF-7 cells alone (MCF-7/DMEM) in Matrigel exhibited the weakest tumorsphere formation capacity (Fig. 6B). RT-qPCR analysis further revealed a higher expression of key CSC markers SOX2 and OCT4 in AM-CM treated MCF-7 (MCF-7/AM-CM) than in the MCF-7 cells alone (MCF-7/DMEM; Fig. 6C and D).

To further evaluate the effect of ADSCs on tumor growth of MCF-7 cells in vivo, a xenograft model we established, in which MCF-7 cells were mixed with or without ADSCs and were then inoculated subcutaneously into nude mice. As shown in Fig. 7A and B, the tumor volume of the ASC-treated group was markedly increased compared with that of the control group. H&E staining revealed that the necrotic area in ASC-treated tumor tissue was considerably reduced compared with that in controls (Fig. 7C and D). These results indicate that ADSCs may enhance the stemness expression and tumor-promoting properties of MCF-7 cells.
Expression of epithelial and mesenchymal markers. In view of the morphological changes of MCF-7 cells in tumorspheres, whether the acquisition of stemness properties is associated with the epithelial to mesenchymal transition (EMT) was investigated. Representative EMT markers were analyzed in the AM-CM treated MCF-7 cells group (MCF-7/AM-CM) and the control MCF-7 cells group (MCF-7/DMEM). Results revealed reduced E-Cadherin (epithelial marker) and increased Vimentin (mesenchymal marker) mRNA expression in the AM-CM treated MCF-7 cells compared with the control MCF-7 cells (Fig. 8A). Protein expression analysis revealed similarly decreased E-Cadherin and increased Vimentin level in AM-CM treated MCF-7 cells compared with control MCF-7 cells (Fig. 8B), indicating that the ADSCs may induce the EMT in MCF-7 cells.

Discussion

Although stem cells are a promising source for cell therapy in regenerative medicine, the potential pro- or anti-tumoral actions of these cells remain controversial (29). Adipose tissue is an abundant, accessible and rich source of ADSCs, and adipose transplantation is gaining increasing interest among the plastic surgery techniques currently available to reconstruct the breast after mastectomy for breast cancer. However, recent scientific attention has turned to whether grafted ADSCs within adipose tissue may increase the risk of cancer recurrence (14, 19, 30-33).

In the present study, ADSCs and MCF-7 cells were co-cultured in a 3D model to investigate the impacts of ADSCs on breast cancer cells. It was found that the co-culture system resulted in migration of ADSCs to MCF-7 cells and simultaneously promoted tumor progression. Similar to other immune cells, MSCs exhibit tropism for sites of tissue damage and the tumor microenvironment (34-36). Various studies indicated that MSCs migrated to sites of inflammation and diseased tissues when injected systemically (37, 38). In contrast, other studies have reported that MSC migration can be induced by conditioned medium from colorectal cancer (39), gliomas (40, 41), and breast cancer (42) cells in vitro. To date, the majority of the studies on MSC tumor tropism were performed with bone marrow-derived MSCs, and limited data are available regarding the ADSCs from adipose tissue. In the present study, it was observed that MCF-7 cells induced efficient ASC tropism. MSC migration to tumors is thought to be due to chemokines secreted by tumor cells, but this needs to be validated. Chemokines were originally identified as potent attractants for leukocytes, such as neutrophils and monocytes, and were generally regarded as mediators of acute and chronic inflammation (inflammatory chemokines) (43). Additionally, chemokines and their receptors have been identified as actors promoting MSC tumor tropism and initiation or cancer progression (43-48). Among these inflammatory chemokines, MIP-1α and MIP-3α are considered as key considered factors in inducing MSC migration (39). MIP-1α and MIP-3α are cytokines that mainly regulate immune cell migration (49) and were recently considered serum biomarkers for hepatocellular carcinoma (50). Lejmi et al (36) investigated the migration of human bone marrow-derived MSCs induced by conditioned medium of Huh-7 hepatoma cells, detecting increased levels of MIP-1α and MIP-3α in Huh-7-CM using a human cytokine antibody array. Transwell migration assay showed that recombinant MIP-1α and MIP-3α increased bone marrow-derived MSC migration and that inhibition of antibodies against...
MIP-1δ and MIP-3α slightly decreased MSCs migration. Consistent with these results, increased expression levels of MIP-1δ and MIP-3α were detected in the co-culture system compared with the control, indicating that these two inflammatory chemokines may participate in ADSCs migration in the present study.
Along with the migration of ADSCs to MCF-7 cells, aggregation and sphere-like structure formation of MCF-7 cells are also important features of the 3D co-culture system. Previous studies have reported that sphere-like structure formation of cancer cells in vitro indicates the acquisition of CSC properties in cancer cell subpopulations (51-53). CSCs are identified by high self-renewal capability, the capacity to grow as tumourspheres in vitro and tumor growth promotion in vivo (54,57). In xenograft breast cancer models, MCF-7 and ADSCs co-injection induced notably faster tumor growth and fewer necrotic areas in tumor tissues than MCF-7 injection alone, suggesting the cancer-promoting activity of ADSCs in vivo. Prompted by these results, it could be assumed that tumoursphere and tumor tissue formation is associated with CSC generation. The CSC markers SOX2 and OCT4 have been reported to inhibit apoptosis and promote biological activity in CSCs (55,56). SOX2 and OCT4 expression levels were increased in MCF-7 cells after treatment with AM-CM on Matrigel substrates. This result indicated that the tumoursphere formation in the co-culture system is likely associated with the stem-like transfer of MCF-7 cells influenced by the secretory cytokines from ADSCs.

Candidate CSCs have been identified in a variety of human malignancies, including leukemias, and a number of solid tumors, such as glioblastomas, medulloblastomas and carcinomas (57-60). The contributions of the EMT program in promoting cancer cells with stem-like properties have been well documented in many types of carcinoma. Mani et al. (61) demonstrated a direct link between the EMT and epithelial stem cell properties. Using different EMT inducers, they showed that the induction of the EMT in human breast cancer cells accounted for the acquisition of their stem-like characteristics. The EMT is a key program during embryonic development, tissue remodeling, and cancer progress (62) and is characterized by the loss of epithelial characteristics coupled with the gain of mesenchymal properties. Furthermore, EMT process is associated with a mesenchymal-like breast cancer phenotype, including acquisition of invasive properties and the loss of cell-cell adhesion. These features are associated with a more aggressive phenotype and a poor prognosis (63). In direct co-culture systems, malignant tumoursphere formation was observed, indicating loss of contact inhibition. Hence, we hypothesized that co-culturing with ADSCs promoted the EMT of MCF-7 cells. As expected, gene expression analysis revealed a significant upregulation of the mesenchymal marker Vimentin and the downregulation of the epithelial marker E-Cadherin in AM-CM treated MCF-7 cells compared with the control, indicating the potential acquisition of EMT process in MCF-7 cells. Altogether, the present results indicated that the stem-like transfer of MCF-7 cells may be a consequence of the EMT induced by secretory cytokines from ADSCs.

Matrigel-based 3D co-culture system may be a convenient and rapid platform for studying cell interaction in tumor development (64,65). Using this platform, it was demonstrated that the interaction between ADSCs and MCF-7 cells stimulated the expression of the chemokines MIP-1δ and MIP-3α, which may act as a regulator inducing the migration activity of ADSCs toward breast cancer cells and to establish direct cell-cell contacts. Furthermore, it was demonstrated that ADSCs serve pro-malignant roles in MCF-7 cells through promoting the tumoursphere formation of MCF-7 cells, which are likely associated with CSC properties through the EMT process (Fig. 9). The promoted tumorigenicity was further confirmed in the in vivo xenotransplantation model. These results are important for safety concerns regarding the clinical application of ASC-based strategies, such as fat grafting, in post-oncologic breast reconstruction, interestingly because microscopic tumor cells may remain after tumor resection.

However, the present study is small and data is limited, several drawbacks should be resolved to fully clarify the precise mechanisms of ADSCs facilitating breast cancer development. First, although the procedure for ADSCs isolation and expansion is considered the most widely used method (66), these cultured cells are still heterogeneous, containing stem cells with different multipotential properties, committed progenitors, and differentiated cells, which in turn may affect the biological properties of the total population. Therefore, using a purified stem cell population from adipose tissue would help improve the study design and strengthen our conclusion. Second, MIP chemokines were not directly shown to act on the migration activity of ADSCs towards MCF-7 cell due to current limited conditions. According to previous studies, MIP cytokines mainly regulate immune cell migration (53) and were recently found to be important chemoattractants that induce MSC migration and further favor its differentiation (36). Using recombinant chemokines MIP-1β and MIP-3α, Lejmi et al (36) revealed that MSC migration was induced whereas addition of anti-MIP-1β and anti-MIP-3α antibodies decreased the MSC migration activity. These results may help to demonstrate that MIP-1β and MIP-3α may be involved in the migration of ADSCs to MCF-7 cells in the present study but using recombinant chemokines MIP-1β and MIP-3α or knockdown of these genes to solidify this conclusion is still necessary in further studies. Third, interaction between the other types of breast cancer cells and ADSCs should be investigated in further studies to fully clarify the potential impacts of fat grafting on breast cancer cells in the clinic.

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Availability of data and materials

The data sets used and/or analyzed during the present study are available from the corresponding author on reasonable request.
Authors’ contributions

YH conceived, designed the study and wrote the manuscript. YC and XW performed the experiments and analyzed the results. JG and FL designed and coordinated the study. All authors read and approved the manuscript and agree to be accountable for all aspects of the research in ensuring that the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Ethics approval and consent to participate

All procedures performed in the present study involving human participants were approved by the Southern Medical University Institutional Review Board (Guangzhou, China) and the patient provided written informed consent to donate remaining tissues after liposuction. All procedures performed involving animal experiments were approved by the Nanfang Hospital animal ethics committee (permit no. NFFY201679) and was conducted in accordance with the ethical standards of the National Health and Medical Research Council China.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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