Therapeutic effects of adipose-derived stem cells-based microtissues on erectile dysfunction in streptozotocin-induced diabetic rats

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This study aimed to explore the therapeutic effects of adipose-derived stem cells (ADSCs)-based microtissues (MTs) on erectile dysfunction (ED) in streptozotocin (STZ)-induced diabetic rats. Fifty-six 8-week-old Sprague-Dawley rats received intraperitoneal injection of STZ (60 mg kg−1), and 8 weeks later, the determined diabetic rats randomly received intracavernous (IC) injection of phosphate buffer solution (PBS), ADSCs, or MTs. Another eight normal rats equally got IC injection of PBS. MTs were generated with a hanging drop method, and the injected cells were tracked in ADSC- and MT-injected rats. Four weeks after the treatments, intracavernous pressure (ICP), histopathological changes in corpus cavernosum (CC), and functional proteins were measured. Rat cytokine antibody array was used to detect ADSCs or MTs lysate. The results showed that MTs expressed vascular endothelial growth factor (VEGF), nerve growth factor (NGF), and tumor necrosis factor-stimulated gene-6 (TSG-6). MTs injection had a higher retention than ADSCs injection and MTs treatment improved ICP, neuronal nitric oxide synthase (nNOS) expression, smooth muscle, and endothelial contents in diabetic rats, ameliorated local inflammation in CC better. Thus, our findings demonstrate that IC injection of MTs improves erectile function and histopathological changes in STZ-induced diabetic rats and appears to be more promising than traditional ADSCs. The underlying mechanisms involve increased cell retention accompanied with neuroprotection and anti-inflammatory behaviors of the paracrine factors.

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INTRODUCTION
Erectile dysfunction (ED) is more prevalent in men with diabetes mellitus (DM). The morbidity rate of ED among diabetic men varies from 35% to 90% due to different population and ages.1,2 Aside from the heavy burden, diabetes mellitus induced erectile dysfunction (DMED), pathogenic manifestations involving endothelial injury, neuropathy, microvascular and fibrous-muscular alterations is usually more serious and difficult to treat than nondiabetic.3,4 The oxidation of low-density lipoproteins and overproduction of oxygen-free radicals induced by hyperglycemia may lead to smooth muscle dysfunction.5 Hyperglycemia can result in mitochondrial fragmentation and apoptosis of endothelial cells.6 Moreover, DMED is highly associated with nerve growth factor (NGF) deficiency and impaired vascular endothelial growth factor (VEGF) signaling.7,8 Phosphodiesterase type 5 inhibitor (PDE5i) was considered to be the first-line pharmacologic therapy for ED since 1998; however, PDE5i showed less efficacy in treating DMED.9

Recently, many experimental approaches for DMED have emerged, including insulin treatment,7 antioxidant therapy,10 low energy shock wave therapy,11 stem cells, and gene therapy.12,13 Among these strategies, stem cell-based therapy is considered to be promising due to its ability to recover functional cells and tissues. For choosing the candidate stem cells, adipose-derived stem cells (ADSCs) have been proposed as one of the most suitable types. Similar to bone marrow stem cells (BMSCs), ADSCs possess the ability of self-renewing and differentiation.14 ADSCs can be collected in abundant quantities and harvested by a minimally invasive procedure. In addition, the successful transplantation of allogeneic ADSCs indicated a low immunogenicity of the cells.15 A growing body of evidence suggested the usage of ADSCs in several ED models.16-18 Of interest, the benefits of ADSCs for ED appear to be related to growth factor and cytokine activity.19 However, the single-cell-based injection is known as low cellular survival which is necessary to long-term success. Adult stem cells might lose many properties in ordinary adherent culture.20 Numerous reports noticed that adult stem cells cultured as spheroids which were scaffold-free could increase the therapeutic potential.18,19 Our previous study showed that ADSCs-based microtissues (MTs) improved erectile function through neuroprotection in a rat model of cavernous nerve injury.20 In
this study, the efficacy of ADSCs and ADSC-based MTs was compared for the treatment of ED in streptozotocin (STZ)-induced diabetic rats and histologic changes were examined.

**MATERIALS AND METHODS**

**Animals**

A total of 64 male Sprague-Dawley rats aged 8 weeks old (body weight: 270–310 g) were purchased from the Animal Center of Peking University. The Committee for Animal Care and Use of Peking University approved the experiments. After harvesting of paratesticular fat, 56 rats were fasted for 16 h, and then intraperitoneally injected with STZ (60 mg kg⁻¹; Sigma-Aldrich, St. Louis, MO, USA). Blood glucose levels were measured at 72 h after STZ injection using a blood glucose meter (B. Braun, Melsungen, Germany). Of the 56 rats, 54 (96.43%) were diabetic with a fasting glucose concentration higher than 300 mg dL⁻¹.

An intraperitoneal injection of phosphate buffer solution (PBS) was administered to 8 rats as control group. At 8 weeks after STZ injection, apomorphine (100 μg kg⁻¹; Sigma-Aldrich, St. Louis, MO, USA) was used to screen the diabetic rats. DMED was present in 48 of the 54 (88.89%) rats. The 48 DMED rats were numbered and randomly assigned to three groups: intracavernous (IC) injection of PBS (n = 8, the DM + PBS group), ADSCs (n = 20, the DM + ADSCs group), and MTs (n = 20, the DM + MTs group). The control group also received an IC injection of PBS. ADSCs were labeled with the chloromethylbenzamido derivative 1,19-dioctadecyl-3,3,39,39-tetramethylindocarbocyanine perchlorate (CM-Dil; Molecular Probes, Carlsbad, CA, USA) and tracked at days 1, 7, and 14 (n = 4) after injection. At day 28, rats in each group were examined for ED before the harvest of tissues.

**ADSCs isolation and MTs generation**

ADSCs were isolated from paratesticular fat and cultured as previously standardized method. In brief, all the animals underwent lower abdominal midline incision and bilateral resection of paratesticular adipose tissue. The adipose tissue was rinsed with PBS containing 1% Streptomycin and Penicillin, chopped into small pieces, and incubated in 0.075% collagenase type IA (Sigma-Aldrich, St. Louis, MO, USA) for 80 min. The top lipid part was removed and the liquid part was centrifuged at 1000 g at room temperature for 10 min. Then, the remaining cells were suspended in low glucose Dulbecco’s modified Eagle’s medium (DMEM, HyClone, Logan, UT, USA) supplemented with 1% Streptomycin and Penicillin and 10% fetal bovine serum (PBS, HyClone, Logan, UT, USA). After filtered through a 100-μm cell strainer, the suspension was plated in a 10-cm dish and cultured at 37°C in 5% CO₂.

MTs were generated with ADSCs or 3-day-old MTs (1 × 10⁶ ADSCs per MT) in 100 μl PBS. After filtered through a 100-μm cell strainer, the suspension was plated in a 10-cm dish and cultured at 37°C in 5% CO₂. MTs were generated with ADSCs or 3-day-old MTs (1 × 10⁶ ADSCs per MT) in 100 μl PBS. When reaching approximately 80% confluence, ADSCs in each dish were centrifuged and resuspended in 1.2 ml DMEM. Then, ADSCs (1 × 10⁶ cells in 30 μl) were dropped onto the cover of new dishes (40 drops per dish), 5 ml PBS was added to each dish, and the cover was carefully placed back on the dish. Dishes were kept at 37°C in 5% CO₂ for 3 days.

**ADSCs and MTs characteristics**

Osteogenic and adipogenic differentiations were performed on both ADSCs and MTs. ADSCs and 3-day-old MTs were seeded in 6-well tissue culture plates and cultured in rat adipose-derived stem cell osteogenic or adipogenic differentiation medium (Cyagen, Santa Clara, CA, USA). We replaced the medium every 3 days with the differentiating time of 21 days, removed the differentiation medium, and rinsed the well with PBS. The cells were fixed with 4% formaldehyde solution for 30 min, and then stained with Alizarin Red or Oil Red O working solution for 30 min. After rinsing, the cells were visualized under light microscope (Leica, Heidelberg, Germany).

Cell lysate from ADSCs or MTs was mixed with biotinylated detection antibodies, and then incubated with a rat cytokine antibody array membrane (R&D systems, Minneapolis, MN, USA) which containing capture antibodies of 29 different target proteins. After washing, the membrane was exposed using chemiluminescent detection reagents.

**IC injection**

Under aseptic conditions, the determined DMED and normal rats were anesthetized with 10% chloral hydrate (300 mg kg⁻¹). The penis was exposed in each group, and a 24-gauge needle was used to inject a total of 1 × 10⁶ ADSCs or 100 MTs (1 × 10⁴ ADSCs per MT) in 100 μl PBS or only 100 μl PBS into the corpus cavernosum (CC). An elastic band was applied to the base of the penis and the pressure was maintained for 2 min after the injection.

**Measurement of erectile function**

Erectile function was determined by intracavernosal pressure (ICP) and mean arterial pressure (MAP) 4 weeks postinjection. Under 10% chloral hydrate, the major pelvic ganglion (MPG) and cavernous nerves (CN) were exposed through a midline laparatomy. The penile was exposed by removing overlying skin and ischiocavernous muscle. One of the 24-gauge needles that were connected to PE-50 tubes with heparinized saline (250 IU ml⁻¹) was inserted into the left carotid to measure MAP. The other one was inserted into corpus cavernosum (CC) to measure ICP. PE-50 tubes were connected to the data acquisition system (MP150, BIOPAC Systems Inc., Goleta, CA, USA). The CNs were stimulated using a stainless steel bipolar hook electrode with the following parameters: 20 Hz, pulse width of 0.2 ms, 1.5 mA, for 50 s. The ratio of maximal ICP (mm Hg) to MAP (mm Hg) was calculated.

**Immunofluorescence staining**

For immunofluorescence (IF) staining, the penile tissue (midshaft portion) and entire major pelvic ganglia (MPG) were fixed in fresh 4% paraformaldehyde, and then were immersed in 30% sucrose in PBS overnight at 4°C. The fixed tissues and 3-day-old MTs were cryosembbed in optimal cutting temperature compound (Sakura Finetek, Torrance, CA, USA) and cut into 5 μm sections before mounting on slides. MPGs were cut crosswise in the middle. After permeabilization and blocking, the slides were incubated with primary antibodies, including rabbit anti-neuronal nitric oxide synthase (nNOS, 1:200; Abcam, Cambridge, UK), rabbit anti-NGF (1:400; Abcam, Cambridge, UK), rabbit anti-VEGF (1:200; Abcam, Cambridge, UK), rabbit anti-α smooth muscle actin (α-SMA, 1:1000; Abcam, Cambridge, UK), rabbit anti-von Willebrand factor (vWF, 1:2000; Abcam, Cambridge, UK), rabbit anti-CD31 (1:400; Abcam, Cambridge, UK), rabbit anti-CD34 (1:200; Abcam, Cambridge, UK), rabbit anti-CD105 (1:400; Abcam, Cambridge, UK), and mouse anti-tumor necrosis factor-induced protein-6 (TSG-6, 1:200; Abcam, Cambridge, UK) at 4°C overnight. At room temperature, the sections were rinsed and incubated with Alexa Fluor 594 conjugated secondary antibodies (Invitrogen, Carlsbad, CA, USA). Nuclei were stained by 4',6-diamidino-2-phenylindole (DAPI, Invitrogen) for 5 min. For tracking the labeled cells, the slides of penile tissues were stained with DAPI only. Slides were visualized under a fluorescence microscope (Leica, Heidelberg, Germany).

**Western blotting**

Total proteins of rat penis, ADSCs, and 3-day-old MTs were extracted by mechanical homogenization in lysis buffer containing protease and phosphatase inhibitors (Sigma-Aldrich, St. Louis, MO, USA).
Proteins were quantified using a bicinchoninic acid (BCA) protein assay kit (Thermo Scientific, Waltham, MA, USA). A volume of 20 μg sample of each protein was electrosaturated in a 10% sodium dodecyl sulfate-polyacrylamide gel and transferred to a polyvinylidenefluoride membrane (Millipore Corporation, Bedford, MA, USA). For incubation, the primary antibodies were rabbit anti-NGF (1:400), rabbit anti-nuclear factor kappa-B (NF-kB, 1:2000), rabbit anti-VEGF (1:200), rabbit anti-α-SMA (1:2000), rabbit anti-nNOS (1:400), mouse anti-tumor necrosis factor-induced protein-6 (TSG-6, 1:200; all from Abcam, Cambridge, UK), and mouse anti-glyceraldehyde phosphate dehydrogenase (GAPDH, 1:10 000; Santa Cruz Biotechnology, Santa Cruz, CA, USA). After incubation with the appropriate secondary antibody (1:2000; Santa Cruz Biotechnology), membranes were exposed using chemiluminescent detection reagents. Images were obtained with a C-digit machine (LI-COR Biosciences, Cambridge, UK).

**Image and statistical analysis**

Histomorphometric results were representative values (n = 5 per group) and analyses for slides were performed using Image-Pro Plus 6.0 software (Media Cybernetics, Bethesda, MD, USA). Western blotting (WB) and rat cytocrine array results were analyzed with ImageJ (National Institutes of Health, Bethesda, MD, USA). Statistical analyses were performed with the SPSS 19.0 (SPSS Inc., Chicago, IL, USA). Results were expressed as mean ± standard deviation (s.d.) and values of P < 0.05 were considered statistically different. Multiple comparisons between groups were done using one-way analysis of variance followed by a post hoc analysis with the Tukey–Kramer test while a comparison between two groups was performed by t-test.

**RESULTS**

**Rats blood glucose and body weight**

One rat in DM + PBS group and two rats in DM + ADSCs group were dead and abandoned during the experiment. The final blood glucose levels of STZ-induced diabetic rats were remarkably higher compared to PBS-treated control rats (P < 0.05). The final weights of diabetic rats were markedly lower than normal rats (P < 0.05). Among DM + PBS, DM + ADSCs, and DM + MTs groups, blood glucose levels and body weights were not statistically different (Table 1).

**Characterization of ADSCs and MTs**

ADSCs cultured in a hanging drop gradually aggregated into a single central spheroid (Figure 1a), and the diameter of spheroid reduced around 175.11 ± 5.58 μm at the third day (Figure 1b). CM-Dil-labeled cells were shown in ADSCs or MTs. Immunofluorescence results showed that CD31 and CD34 were negative in MTs while CD105, vWF, VEGF, NGF, and TSG-6 were expressed in MTs (Figure 1c). Osteogenic and adipogenic differentiations demonstrated that both ADSCs and MTs have the potential for multipotency (Figure 1d). The endothelium content was indicated by vWF-positive area in Figure 1c. The endothelium content was indicated by vWF-positive area in Figure 1d.

**The disappearance of labeled cells in ADSCs- and MT-injected rats**

The CM-Dil-labeled cells were tracked in ADSCs- and MT-injected rats during days 1, 7, and 14 post-IC injection (Figure 2a). As they were showed by integral optical density (IOD), cells disappeared rapidly in ADSC-injected rats compared with MT-injected rats (Figure 2b). At days 7 and 14, cells remained retaining in MT-injected rats while very few cells could be observed in ADSC-injected rats (P < 0.05).

**Erectile function assessment**

Diabetic rats in DM + PBS group showed significant decreases in both maximum ICP and ICP to MAP ratios compared to the normal rats (P < 0.05). ADSCs treatment partially ameliorated erectile dysfunction compared with PBS-treated diabetic rats (P < 0.05) (Figure 3a) while MT-injected rats displayed significant improvements compared with ADSC-injected rats (P < 0.05). There were no striking differences in MAP between groups (Figure 3a and 3b).

**Smooth muscle and endothelium contents in cavernous tissue**

If staining and WB analysis of α-SMA expression in penis showed a significant decrease of smooth muscle content in PBS-treated diabetic rats (Figure 4). Both ADSCs and MTs treatments restored the smooth muscle content (P < 0.05). Surprisingly, MTs treatment displayed a better recovery with the higher smooth muscle density (P < 0.05). The endothelium content was indicated by vWF-positive area in Figure 4.

**Table 1: Body weight and blood glucose variables**

| Variable                  | Control       | DM + PBS       | DM + ADSCs     | DM + MTs       |
|---------------------------|---------------|----------------|----------------|----------------|
| Initial weight (g)        | 300.72±11.32  | 299.02±9.91    | 300.22±10.02   | 302.08±10.72   |
| Final weight (g)          | 605.84±43.82  | 370.03±23.70*  | 387.06±19.72*  | 395.20±29.58*  |
| Initial glucose (mmol l⁻¹) | 6.52±0.60     | 6.46±0.71      | 6.58±0.80      | 6.92±0.48      |
| Final glucose (mmol l⁻¹)  | 6.62±0.52     | 27.82±5.12*    | 27.66±2.72*    | 29.80±3.81*    |

Values are expressed as means±s.d. *P<0.05 when compared with control group. s.d.: standard deviation; PBS: phosphate buffer solution; DM: diabetes mellitus; ADSCs: adipose-derived stem cells; MTs: micratissues.
CC and markedly decreased in rats of DM + PBS group compared with rats in control group \( P < 0.05 \). After IC injection of ADSCs, the endothelium content was increased to some degree \( P < 0.05 \). After MTs treatment, the endothelium in cavernous tissue was better restored compared with ADSC-injected rats \( P < 0.05 \) (Figure 4a and 4c). Meanwhile, WB results showed that VEGF expression in penis was higher in MT-injected rats than that of ADSC-injected rats \( P < 0.05 \) or PBS-injected diabetic rats \( P < 0.05 \) (Figure 4d and 4f).

**DISCUSSION**

Nitricergic nerves, endothelium, and smooth muscles are crucial for normal penile erection. The released nitric oxide (NO) from endothelium or nerve terminals stimulates the production of cyclic guanosine monophosphate (cGMP), and then the gradually accumulated cGMP helps dilate corporal smooth muscle, facilitate blood supply, and cause erection. Macrophasculopathy, microvasculopathy, and endothelial dysfunction are the major vasculopathy limiting the blood flow in the penis. Microvascular disease in diabetes also causes nerve ischemia resulting in peripheral neuropathy and autonomic neuropathy which play important roles in DMED. In the present study, reductions of smooth muscle and endothelium density in CC were observed in PBS-injected diabetic rats. Moreover, diabetic rats showed neuronal degeneration through markedly reduced nNOS content in DPN and nNOS-positive neurons in MPG. These results are consistent with previous findings about diabetic rats. As for ED therapy, independent groups have investigated the feasibilities and advantages of ADSCs in rat models of DMED, cavernous nerve injury, and Peyronie’s disease. These results were encouraging but the traditional single-cell-based injection was relative preliminary, and the therapeutic

**Figure 2:** Time-dependent disappearance of IC-injected cells. (a) Representative images of CM-Dil-labeled cells retained in corpus cavernosum at days 1, 7, and 14 postinjection in ADSC- and MT-injected rats. Original magnification: Left, ×200 and right, ×1000. (b) Quantitative data of labeled cells between two groups are expressed as integrated optical density (IOD) of CM-Dil-positive area in corpus cavernosum. \( * P < 0.05 \) compared with DM + ADSCs group.

**Figure 3:** Erectile function of each group. (a) Representative images of intracavernous pressure (ICP) and mean arterial pressure (MAP). Horizontal bar below ICP represents 50 s of stimulate duration. (b) Left: Results of peak ICP increase of ADSCs- and MTs-treated rats. Right: Results of ratio ICP to mean MAP of each group. \( * P < 0.01 \) when compared with DM + PBS group. \( \ddagger P < 0.05 \) when compared with DM + ADSCs group.
potential was hampered by poor retention rate due to the blood flow. In this regard, MTs generated by a hanging drop method was seen as superior to single cell strategy. To examine whether IC injection of MTs has a better retention in diabetic rats, we labeled ADSCs with CM-Dil.
before treatment. The results showed that the labeled cells disappeared rapidly in CC of ADSC-injected rats as very limited amount was detected 2 weeks after IC injection while more CM-Dil-positive cells could be observed in MT-injected rats. These are consistent with the previous studies that rare cells could be tracked in cavernous tissue after transplantation and stem cells preprocessed as MTs could promote the cell retention and engraftment. The higher local retention could change the extracellular compartment of local tissue and might contribute to the greater improvement of erectile function.

Besides, many works have proposed that the transplanted cells played a part in CC might not through direct differentiation into local cell types but through the secretion of cytokines and growth factors. In this study, WB results demonstrated that the expressions of VEGF and NGF were higher in MTs than those of ADSCs. As a proangiogenic factor, VEGF has multiple functions including stimulates proliferation, inhibits apoptosis, and promotes cell survival. Impaired VEGF signaling pathway was closely related to endothelial dysfunction and was thought to be a possible cause for ED in diabetes. NGF is a type of neurotrophins that promote neural survival and differentiation. It has been revealed that NGF expression in target organs was decreased and the axonal transport of NGF was delayed in diabetic rats. VEGF and NGF expressions in penis were better augmented in MT-injected rats than those of ADSC-injected rats. Smooth muscle and endothelium density accompanied with nNOS content were increased more in MTs-treated rats than those of ADSCs-treated rats. These results demonstrated that MTs injection had better effects on vascularization and neuroprotection.

ADSCs expressed higher levels of the inflammatory proteins CINC-1, CCL5, and IL-1ra than MTs, indicating that MTs have anti-inflammatory activity. Indeed, MTs, but not ADSCs, expressed TSG-6, a multifunctional protein with an anti-inflammatory effect. Importantly, TSG-6 induced anti-inflammatory activity through the inhibition of transcript factor NF-κB, which regulated many key genes related to immune and inflammatory responses. WB results also showed the lower NF-κB expression in MTs compared with ADSCs, all of these indicated the anti-inflammatory ability of MTs. To examine whether IC injection of ADSCs or MTs inhibits local inflammation, we detected NF-κB expression in penis of each group. Interestingly, WB results demonstrated that diabetes-induced inflammation in penis and ADSCs treatment did not affect the local inflammation while the inflammatory state in CC was partially recovered by IC injection of MTs. This could be another reason for the higher efficiency of MTs treatment. Further studies should be carried out to check the systemic inflammation by measuring both pro- and anti-inflammatory cytokines to better understand the anti-inflammatory function of MTs.

Culturing adult stem cells in adherent monolayer could not truly recreate the specialized microenvironment which was known as stem cell niche in vivo and might limit the potential of transplanted cells. Therefore, mimicking the niche composition might help enhance the regenerative potential of ADSCs expanding in vitro. Moreover, subculturing of stem cells upon long-term in vitro would lead to phenotypes change and some abilities lose. In this study, both single ADSCs and ADSCs dissociated from MTs possessed the adipogenic and osteogenic differentiation potential indicating that ADSCs retained their properties when cultured as MTs. Similarly, previous work on mesenchymal stem cells demonstrated that cells in spheroids retained most of the surface epitopes and differentiation.
potential. Nutrient deprivation and "microgravity" are the possible stimulating factors related to the changes of cells cultured as MTs. It is possible that stem cells cultured in MTs can contact closely with each other and the large amount of the cells ensures the quality of cell signaling and secretion. This can be partly supported by the higher expression of ICAM-1 in MTs since ICAM-1 was important in stabilizing cell-cell interactions.  

IC injection of ADSCs-based MTs resulted in a better improvement of erectile function in diabetic rats than single-cell ADSCs injection, and the greater therapeutic effects through enhanced paracrine abilities were supported by indirect evidence in our study. Moreover, the erectile function and histologic changes associated were not completely recovered only by ADSCs or MTs treatment. The diabetes status is relative complex and the hyperglycemia may cause sustained damages; therefore, blood glucose controlled by insulin combined with MTs injection will be a more promising way for DMED treatment. Investigations should be carried out in more ED animal models and to better evaluate the therapeutic effects and safety for potential clinical application.

AUTHOR CONTRIBUTIONS  
JQH, ZCX, FZ, and YH conceived and designed the experiments. FZ, YH, YDX, HEL, RLG, and HX performed the experiments. BCY and JQH, ZCX, FZ, and YH involved in the retrieval and analysis of the data. YH, FZ, JQH, and ZCX participated in the drafting and final editing. All authors read and approved the final manuscript.

COMPETING INTERESTS  
All authors declared no competing interests.

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