Mesenchymal stem cell-derived small extracellular vesicles mitigate oxidative stress-induced senescence in endothelial cells via regulation of miR-146a/Src

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Senescent endothelial cells (ECs) could impair the integrity of the blood vessel endothelium, leading to vascular aging and a series of diseases, such as atherosclerosis, diabetes. Preventing or mitigating EC senescence might serve as a promising therapeutic paradigm for these diseases. Recent studies showed that small extracellular vesicles (sEV) have the potential to transfer bioactive molecules into recipient cells and induce phenotypic changes. Since mesenchymal stem cells (MSCs) have long been postulated as an important source cell in regenerative medicine, herein we investigated the role and mechanism of MSC-derived sEV (MSC-sEV) on EC senescence. In vitro results showed that MSC-sEV reduced senescent biomarkers, decreased senescence-associated secretory phenotype (SASP), rescued angiogenesis, migration and other dysfunctions in senescent EC induced by oxidative stress. In the in vivo natural aging and type-2 diabetes mouse wound-healing models (both of which have senescent ECs), MSC-sEV promoted wound closure and new blood vessel formation. Mechanically, miRNA microarray showed that miR-146a was highly expressed in MSC-sEV and also upregulated in EC after MSC-sEV treatment. miR-146a inhibitors abolished the stimulatory effects of MSC-sEV on senescence. Moreover, we found miR-146a could suppress Src phosphorylation and downstream targets VE-cadherin and Caveolin-1. Collectively, our data indicate that MSC-sEV mitigated endothelial cell senescence and stimulate angiogenesis through miR-146a/Src.

INTRODUCTION
Aging induced vascular dysfunctions play a critical role in the pathogenesis of a variety of age-related diseases, such as delayed wound healing, heart failure, diabetes, Alzheimer’s disease, and kidney diseases1,2. In demographic research, vascular diseases are the leading causes of severe long-term impairment and mortality in older adults3,4. Therapeutic strategies to reduce vascular diseases by preventing or eliminating vascular senescence are still required, despite years of intensive research5. Senescent endothelial cells (ECs) exhibit major changes in gene expression, cell replication and morphology, impair the integrity of the endothelium in vascular by influencing the endothelium’s regenerative and angiogenic capacities, reactivity and pathogenic progression, contributing to vascular aging diseases5,7. A range of factors can cause endothelial senescence; among them, oxidative stress has a major role6. H2O2 and high glucose are factors contributing to EC senescence as oxidative stresses9,10. Ming-Hui Zou showed that H2O2 could trigger HUVECs and human aortic smooth muscle cells senescence through Oct4A upregulation11. H2O2-induced EC senescence provokes a DNA-damage response, which results in activation of p53 and p16, the important cell cycle regulatory pathways12. High-glucose induced HUVECs senescence through decreasing the expression level of SIRT3, and increased SA-gal expression and damaged the tube forming ability9.

Oxidative stress participates in the pathogenesis of vascular abnormalities in diabetes which can induce premature senescence via DNA damage, and Streptozotocin(STZ)-induced diabetes can cause ECs senescence13. However, the potential interventions to prevent ECs senescence for the prevention of vascular pathologies are unclear.

Mesenchymal stem cells (MSCs), identified as multipotent stromal cells, have multi-lineage differentiation ability and immunosuppressive properties14 (Supplementary Fig. 1B–F). They can be derived from several sources, including the umbilical cord, bone marrow, or fat tissue, making them ideal for immunomodulation and regeneration as a promising candidate cell type15. Increasing evidence indicates that these features of MSCs are linked to their paracrine activity and extracellular vehicles (EVs) secretion16. Small extracellular vesicles (sEV) are extracellular vesicles generated by fusion with the cellular membrane of multivesicular bodies. They are between 30–150 nm in diameter and contain abundant functional components such as proteins and microRNAs (miRNAs)17. sEV have emerged as a complex means for a variety of cellular processes to be modulated18. sEV derived from MSCs have previously been documented to elicit similar therapeutic effects to their parent MSCs19. More specifically, sEV released by MSCs, such as low immunogenicity, easy storage, and high biosafety, have striking advantages over
whole-cell therapy, sEV derived from MSC have been shown to have great promise in anti-inflammation and injury repair and are commonly studied as a nanotherapeutic agent for stroke and wound-healing treatment. Thus, for certain diseases, MSC-sEV have been described as highly promising cell-free therapeutic agents. Nonetheless, the curative effects and mechanisms of the action of MSC-sEV on senescence are poorly understood, to the best of our knowledge, particularly on EC senescence.

We intended to investigate the effect of MSC-sEV on oxidative stress-induced HUVECs senescence in vitro, and naturally aged and diabetic mice wound-healing model in vivo. Moreover, we investigated the underlying molecular mechanism by using miRNA sequencing and phospho-kinase antibody array. Our results suggested that MSC-sEV can act as a nanotherapeutic agent via mir-146/Sc pathway to prevent oxidative stress-induced EC senescence.

RESULTS

Characterization of MSC-sEV and their active internalization by HUVECs

sEV were isolated from supernatant of MSCs using ultracentrifugation and characterized by western blot, transmission electron microscopy (TEM) and nanoparticle tracking analysis (NTA). sEV-associated protein markers Alix and CD63 (transmembrane/lipid-bound protein) and TSG101 (cytosolic protein) were enriched in MSC-sEV and the negative protein marker calnexin (an endoplasmic reticulum marker) was not found in MSC-sEV compared to MSC-lysate (Fig. 1a). Under TEM, MSC-sEV exhibited classic cup-shaped or sphere-shaped morphology (Fig. 1b). According to NTA, the distribution curve of the particle size of MSC-sEV was between 55 and 200 nm (Fig. 1c). To investigate whether MSC-sEV could be internalized by ECs, we used HUVECs as a cellular model. Supplementary Fig. 1a showed immunofluorescence staining of VWF and CD31 (two typical endothelial markers) in HUVECs. MSC-sEV were labeled with PKH26 and added to HUVECs at a final concentration of 200 ng/μL in the medium. sEV incorporation was observed 2 h after treatment and accumulated in HUVECs over time (Fig. 1e). It has been reported that EVs could be taken up via a variety of endocytic pathways, including macropinocytosis, CME, caveolin-mediated endocytosis, and clathrin- and caveolin-independent endocytosis. A schematic illustration of the different types of endocytosis and their respective inhibitor. Treatment with an inhibitor did not reduce the internalization of PKH26-labeled sEV (Fig. 1e, Supplementary Fig. 1g, h), suggesting that MSC-sEV may gain entry into HUVECs via more than one route. Collectively, we characterized MSC-sEV and showed that they are actively incorporated in vitro by HUVECs via multiple routes.

MSC-sEV mitigated senescence in HUVECs in vitro

H2O2 has been reported to trigger premature senescence by increasing oxidative stress. To investigate the effect of MSC-sEV on senescence, we first established a proper H2O2-induced HUVEC senescence model. HUVECs were exposed to different concentrations of H2O2 (25 μM, 50 μM, 75 μM, 100 μM, 200 μM) for 2 h and several well-established senescence biomarkers were analyzed. H2O2 exposure increased the percentage of senescence-associated β-galactosidase- (SA β-gal) positive cells with senescence-related morphological transformations: enlarged and flat appearance and western blot (Fig. 2d) showed that H2O2 markedly elevated senescence markers p16, p21 and p53, and significantly decreased nuclear morphology marker LaminB1. The effect of H2O2 on HUVECs senescence seemed to be concentration dependent; however, when exceeding 50 μM, the number of adherent cells reduced. Therefore, we chose 50 μM H2O2 in the following experiments. To investigate whether senescence affects uptake of MSC-sEV, we compared the intensity of PKH26 fluorescence in control and senescent HUVECs. No significant differences were observed (Fig. 2e, Supplementary Fig. 1f). These results suggest that senescent HUVECs remained the capacity to uptake MSC-sEV.

Senescent HUVECs were then treated with different concentrations of MSC-sEV (25, 50, 100, 200 ng/μL). MSC-sEV decreased the percentage of SA β-gal positive cells and recovered senescence-related morphological transformations (Fig. 2f, g). qRT-PCR and western blot showed that MSC-sEV markedly reduced the oxidant-induced elevation of senescence markers p16, p21, p53 and significantly increased nuclear morphology marker LaminB1 at both mRNA (Fig. 2h) and protein level (Fig. 2i). When used at 200 ng/μL, MSC-sEV almost completely recovered HUVEC from senescent state in terms of p16, p21, p53, and LaminB1 expression.

High glucose has also been reported to exerted a pivotal role in senescence through oxidative stress. To confirm the senescence reducing effects of MSC-sEV, we induced HUVECs senescence by high glucose and found similar protective effects of MSC-sEV (Fig. 2–l).

We then examined the effects of MSC-sEV on HUVECs functions. MSC-sEV remarkably increased senescent HUVEC migration as evidenced by the transwell migration assay (Fig. 3a, b) and scratch wound assay (Fig. 3c, d). Senescent HUVECs treated with MSC-sEV recovered tube formation in vitro (Fig. 3e, f) and blood vessel formation in Matrigel plugs in vivo (Fig. 3m, n). We also tested proinflammatory cytokines IL-1α, IL-6, and IL-8, which are markers of the senescence-associated secretory phenotype (SASP) and found MSC-sEV suppressed IL-6 and IL-8 expression in senescent HUVECs (Fig. 3g). Additionally, we examined mitochondrial respiration and reactive oxygen species (ROS) levels. Senescent HUVECs had reduced basal mitochondrial O2 consumption rate (OCR) and mitochondrial respiration capacity, and increased ROS levels, which were significantly rescued by MSC-sEV (Fig. 3h, j, k). Moreover, MSC-sEV promoted senescent HUVECs proliferation (Fig. 3l). Taken together, these results suggest that MSC-sEV could prevent oxidative stress-induced senescence in HUVECs.

MSC-sEV promoted wound closure and blood vessel formation in natural aging and type-2 diabetes mice models

The preventive effects of MSC-sEV on oxidative stress-induced senescence in vitro prompt us to investigate their role in wound-healing in vivo. To identify differences between normal and aged mice, we performed wound-healing surgery on 8-week-old mice and 64-week-old mice, and the results indicated that aged mice had a weaker ability of wound healing than younger mice (Supplementary Fig. 2a–c). And we subcutaneously injected PKH26-marked MSC-sEV to the mice dorsal skin, and found that MSC-sEV could be internalized by CD31-positive EC (Supplementary Fig. 2d). We used natural aging mice. Injection of 200 μg MSC-sEV significantly increased wound closure in aged mice as showed by smaller wound areas (Fig. 4a, b). H&E staining was carried out to evaluate the extent of re-epithelialization and scar formation. As shown in Fig. 4c, much wider newly formed epidermis and dermis with hair follicles and fat cells were observed in the wounds treated with MSC-sEV, as compared with the PBS-treated wounds at day 12. Quantitative measurements confirmed that MSC-sEV-treated wounds had a higher rate of re-epithelialization and a lower level of scar formation than the control group (Fig. 4d).

Furthermore, MSC-exosomes-treated wounds exhibited significantly more newly formed blood vessels compared with the control wounds at day 12 post-wounding as demonstrated by immunostaining for endothelial marker CD31 and marker VE-cadherin (Fig. 4e, Supplementary Fig. 2e). Quantitative analysis of the density of new blood vessels confirmed that the beneficial effect of MSC-sEV on revascularization of the wounds (Fig. 4f, Supplementary Fig. 2f).
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Fig. 1 Characterization of MSC-sEV and their active internalization by HUVECs. a Representative images of western blot analysis showing the biomarkers of sEV, including TSG101, CD63, and Alix. Calnexin was used as a negative control (MSC-sEV1, 2.1 mg protein/mL MSC-sEV; MSC-sEV2, 0.21 mg protein/mL MSC-sEV). b Transmission electron microscopic images of MSC-sEV (scale bar, 100 nm). c Nanoparticles tracking analysis reveals the particle distribution of exosomes in various sizes. Within every nanometer diameter sets, the value of the ordinate represents the mean particle number. d Schematic illustration of the different types of endocytosis and their respective inhibitor (Created with BioRender.com). e Uptake of MSC-sEV by HUVEC pretreated with signal inhibitors chlorpromazine (CPZ), nystatin, or 5-(N-ethyl-N-isopropyl)amiloride (EIPA) for 1 h. Cells were incubated with the indicated inhibitors for 1 h before and during the incubation with MSC-sEV. HUVECs incubated with 200 ng/μL PKH26-labeled MSC-sEV for the indicated times, and uptake of MSC-sEV was determined by fluorescence microscopy (scale bar, 100 μm)
Fig. 2 MSC-sEV mitigated senescence in HUVECs in vitro. Cultures exposed to H₂O₂ (25 μM, 50 μM, 75 μM, 100 μM, 200 μM, 2 h) or PBS (control) as showing in the figure. 

- **a** Representative images of senescence-associated β-galactosidase (SA-β-gal) staining (scale bar, 50 μm).
- **b** Quantitation of SA-β-gal-positive cells in HUVEC.
- **c** Quantitative real-time PCR (qRT-PCR) of the senescence markers P16, P21, P53, and LMNB1 mRNA in HUVEC after 24 h exposure to H₂O₂. 
- **d** Representative images of western blot analysis showing the senescence markers P16, P21, P53, and LMNB1 in HUVEC after 48 h of exposure to H₂O₂.
- **e** Representative images of MSC-sEV uptake by HUVEC pretreated with or without H₂O₂ (50 μM, 2 h). HUVECs incubated with 200 ng/μL PKH26-labeled MSC-sEV for the indicated times, and uptake of MSC-sEV was determined by fluorescence microscopy (scale bar, 100 μm) and fluorescence confocal microscopy (scale bar, 20 μm). Next, HUVECs were incubated with 0 ng/μL, 25 ng/μL, 50 ng/μL, 100 ng/μL and 200 ng/μL MSC-sEV for 24 h (c) and 48 h (a, d) after pretreated with H₂O₂ (50 μM, 2 h).
- **f** Representative images of SA-β-gal staining in HUVECs (scale bar, 50 μm).
- **g** Quantitation of SA-β-gal-positive cells in HUVEC. 
- **h** qRT-PCR of the senescence markers P16, P21, P53, and LMNB1 mRNA in HUVEC. 
- **i** Representative images of western blot analysis showing the change of senescence markers P16, P21, P53, and LMNB1 in HUVECs.
- **j** Representative images of SA-β-gal staining in high-glucose-induced senescent HUVECs (scale bar, 200 μm).
- **k** Quantitation of SA-β-gal-positive cells in high-glucose-induced senescent HUVEC.
- **l** Representative images of western blot analysis showing the change of senescence markers P16, P21, P53, and LMNB1 in high-glucose-induced senescent HUVEC.
MSC-sEV rescued senescent HUVECs functions. HUVECs were incubated with (s + 200 ng/μL) or without (s + 0 ng/μL) MSC-sEV for 48 h after pretreated with H₂O₂ (50 μM, 2 h) compared with normal HUVEC (control). a Representative images of transwell migration assays of HUVECs (scale bar, 200 μm). b Quantitation of transwell migration assays of HUVECs. n = 3, **p < 0.01, ***p < 0.001. c Representative images wound-healing assay showing the migratory abilities of HUVECs and the image were taken at the indicated times (scale bar, 200 μm). d Quantitation of wound-healing assay at 12 h. n = 3, ***p < 0.001. e Representative images of in vitro angiogenesis assay (scale bar, 200 μm). f Quantitation of mean tube lengths and branching points in the HUVEC network. n = 3, **p < 0.01, ***p < 0.001. g Quantitation of IL-1 alpha, IL-6, and IL-8 released by HUVEC was detected through ELISA. n = 3, *p < 0.05, **p < 0.01, ***p < 0.001. h Mitochondrial respiratory capacity measured by O₂ consumption rate (OCR) using a Seahorse analyzer. i Quantitation of basal OCR. n = 3, ***p < 0.001. j Quantitation of respiration capacity. n = 3, **p < 0.01. k Quantitation of the ROS level of HUVEC measured by fluorometric intracellular ROS Kit on a flow cytometer. n = 3, **p < 0.01. l Quantitation of HUVEC proliferation evaluated by MTS kits with the OD value on Day 0, Day 1, Day 2, Day 3, and Day 4. n = 4, **p < 0.01, ***p < 0.001. m Representative images of hematoxylin staining and CD31 (red) and DAPI (blue) immunofluorescence staining in paraffin-embedded sections of Matrigel plugs. n Quantitation of tubes formed in vivo. Arrows indicate tubes formed in vivo. n = 3, **p < 0.01.
Fig. 4  MSC-sEV promoted wound closure and blood vessel formation in natural aging and type-2 diabetes mice models. a Representative cutaneous wound photographs in senescent C57BL/6 mice model at days 0, 3, 6, 9, and 12 after the operation treated with PBS (control), MSC-sEV 200 μg around wounds (scale bar, 5 mm). b Quantification of the wound areas during the wound-healing process. n = 6, ***p < 0.001 at Day 12. c H&E staining of the skin tissue around the wound at day 12 after the operation (horizontal arrows indicated the scar width; scale bar, 1 mm). d Quantification of the scar width and percentage of re-epithelialization around the wound. n = 6, **p < 0.01. e Representative immunofluorescence staining images positive cells of CD31 (green staining) and DAPI in paraffin-embedded sections of senescent C57BL/6 mice dorsal skin injected PBS (control) or MSC-sEV. Red arrows indicate tubes formed around the wound (scale bar, 200 μm). f Quantification of vessels formed around the scar. n = 3, ***p < 0.001. g Representative dorsal wound photographs in type-2 diabetes C57BL/6 mice model at day 0, 3, 6, 9, and 12 after the operation treated with PBS (control), MSC-sEV 200 μg around wounds (scale bar, 5 mm). h Quantification of the wound areas during the wound-healing process. n = 6, ***p < 0.001 at Day 12. i H&E staining of the skin tissue around the wound at day 12 after the operation (horizontal arrows indicated the scar width; scale bar, 1 mm). j Quantification of the scar width and percentage of re-epithelialization around the wound. n = 6, **p < 0.01. k Representative immunofluorescence staining images positive cells of CD31 (green staining) and DAPI in paraffin-embedded sections of diabetic C57BL/6 mice dorsal skin injected PBS (control) or MSC-sEV. Red arrows indicate tubes formed around the wound (scale bar, 200 μm). l Quantification of vessels formed around the scar. n = 3, **p < 0.01.
We further examined the functional role of MSC-sEV using a type 2 diabetes mouse model, in which EC senescence also play an important role in its pathogenesis. Diabetic mice were established by injection of STZ and the digital photographs of wounds showed much faster-wound closure in diabetic mice treated with MSC-sEV (Fig. 4g, h). H&E staining showed the extent of re-epithelialization and scar formation in the wounds treated with MSC-sEV was much better than PBS-treated wounds at day 12 after operation (Fig. 4i). Quantitative measurements confirmed that MSC-sEV-treated wounds had a higher rate of re-epithelialization and a lower level of scar formation than the control group (Fig. 4j). Skin images from the undersurface revealed that MSC-sEV-treated wounds exhibited much more newly formed blood vessels compared with the control wounds at day 12 post-wounding (Fig. 4k, Supplementary Fig. 2g). Quantitative analysis of the density of new blood vessels confirmed that the beneficial effect of MSC-sEV on revascularization of the diabetic mice’s wounds (Fig. 4l, Supplementary Fig. 2h). Also, we performed immunofluorescence staining of SA-β gal and CD31 on aged mice and diabetes mice injected with PBS or MSC-sEV, the results showed that MSC-SEV decreased senescent EC number (Supplementary Fig. i–l). Taken together, our data suggested that MSC-sEV promoted skin recovery and angiogenesis in both natural aging and diabetic mouse models.
MSC-sEV rescued senescent HUVEC through up-regulating miR-146a.

As a crucial component of exosome cargo, miRNAs have been reported to play important roles in mediating exosome functions. To identify which miRNA in MSC-sEV contributed to HUVECs senescence, we performed miRNA sequencing of MSC-sEV. Overall, we detected 1038 mature miRNAs, of which 382 had read counts over 100 and we listed the top 20 highly expressed in Fig. 5a.

We then examined changes of miRNAs profile in senescent HUVECs after treatment with MSC-sEV and identified 875 upregulated and 645 downregulated miRNAs (Fig. 5b), among which 63 miRNAs with twofold increase after MSC-sEV treatment were selected (Fig. 5c). We postulated that MSCs exosomal miRNAs contributed to the upregulation of some of the 63 miRNAs, so we overlapped the top 200 miRNAs in MSC-sEV with the 63 miRNAs and found four miRNAs, hsa-miR-146a-5p, hsa-miR-34b-3p, hsa-miR-28-3p, and hsa-miR-412-5p. qPCR analysis confirmed that the four miRNAs were markedly increased in senescent HUVECs after exposure to MSC-sEV (Fig. 5d). We chose hsa-miR-146a-5p and hsa-miR-34b-3p for further analysis as these two miRNAs have been reported to be related to cell senescence. We postulated that if the effects of MSC-sEV were mediated through hsa-miR-146a-5p and has-miR-34b-3p, inhibition of their expression could abolish MSC-sEV effects. So, we transfected senescent HUVECs with respective inhibitors to reduce miR-146a (inhibitor-146a) and miR-34b (inhibitor-34b) before exposure to MSC-sEV. Our data showed that inhibitor-146a but not inhibitor-34b abolished MSC-sEV effects. After miR-146a inhibition, MSC-sEV could not reduce the percentage of SA-β-gal positive cells (Fig. 5e, f), or downregulate p16, p21, p53, or upregulate LMMNB1 expression levels (Fig. 5g, Supplement Fig. 3a). Moreover, after miR-146a inhibition, senescent HUVECs functions including migration and tube formation couldn't be rescued by MSC-sEV (Fig. 5h–k). Whereas, inhibitor-34b had no effects (Fig. 5e–k). These results suggested that MSC-sEV effect was mediated through miR-146a, which prompted us to investigate whether miR-146a could exert preventive effect on senescent HUVEC. We increased miR-146a levels by using mimics of miR-146a (Supplementary Fig. 2b). As expected, mimic-146a decreased the percentage of SA-β-gal positive cells (Fig. 5l, m), downregulated p16, p21, p53, and upregulated LMMNB1 from both mRNA and protein level (Fig. 5n, Supplement Fig. 2c), increased migration and angiogenic capacity (Fig. 5o, p). Collectively, MSC-sEV rescued senescent HUVEC through up-regulating miR-146a.

DISCUSSION

This study was aimed to investigate the therapeutic effects and mechanism of MSC-sEV on oxidative stress-induced HUVECs senescence. We found that in senescent HUVECs, the MSC-sEV treatment protected against senescence-induced biomarkers and dysfunctions, improved angiogenesis, migration, and proliferation capacities, SASP, mitochondrial dysfunction, and ROS level. Furthermore, our data indicated that MSC-sEV could promote skin wound healing in aging and type-2 diabetes mice model after subcutaneous injection of MSC-sEV, suggesting that MSC-sEV may be a potential nanotherapeutic agent for the treatment of vascular-related diseases or disorders, the pathological basis of which involves senescence (Fig. 7).

The use of the traits, included cell function alternations, morphology, gene expression, and positive staining for SA-β-gal.
in combination is the current best practice for identifying senescent cells\(^5\). Thus, we established a convective HUVECs senescence model and found out the concentration of MSC-sEV to rescue senescent HUVECs efficiently in vitro through comprehensive traits mentioned above in this research. EVs derived from stem cells play a key role in mediating the capacity of stem cells through the delivery of biologically active molecules into recipient cells\(^{19}\). Several studies have shown the effects of stem cell-derived EVs in disease therapy\(^{16,21,23,29}\). Antonio Casado-Díaz et al. demonstrated that MSC-derived EV is beneficial to the chronic skin ulcers treatment, including decreasing scar formation and accelerating healing\(^{16}\). MSC-sEV can be in the treatment of neurological diseases such as epilepsy, stroke, or traumatic brain injury which derived from strong anti-inflammatory effects\(^{20}\). Adipose MSC sEV through Wnt/β-catenin signaling improve cell proliferation, migration, and restrain cell apoptosis in skin wound healing\(^{23}\). However, no study reported the effects and underlying mechanisms of MSC-sEV action with respect to ECs senescence. In this research, we first extensively tested some important functions to estimate whether MSC-sEV promote the functional recovery of senescent ECs following identifying senescent biomarkers in HUVECs. EC senescence can impair the repair capacity of the endothelial lining as a direct result of permanent growth arrest\(^{30}\). An in vitro migration and tube formation assay showed that
MSC-sEV could improve the functional recovery of senescent HUVECs compared with the group without sEV. Although the cell proliferation capacity enhanced by MSC-sEV may affect the rate of gap-filling of the scratch wound, the effect of HUVECs on the migration of HUVECs was demonstrated in the transwell migration assay. These results indicate that MSC-sEV contain key factors to restore the proliferation and migration capacity of senescent HUVECs. As the senescence-associated secretory phenotype (SASP), for what is known, senescent cells generate proinflammatory and matrix-degrading molecules, and the SASP discovery proposed a mechanism by which senescent cells could influence tissue and organ function out of proportion to their numbers. Eliminating senescent cells and attenuating the SASP have emerged as promising therapeutic strategies due to their possible role in many aging and disease processes. ELISA tests showed in Fig. 3g suggested that MSC-sEV reduced the increase in SASP subcellular levels. Our results showed that MSC-sEV treatment in induced by senescence (IL-1α, IL-6, and IL-8). Substantial migration of HUVECs was demonstrated in the transwell migration assay showing the phosphorylation ofSrc, VE-cadherin, and Caveolin (downstream of Src) in high-glucose-induced senescent HUVEC. HUVEC was co-cultured with MSC-sEV (200 ng/μl, 4 h) after treated with high glucose (30 mM, 48 h). Representative immunofluorescence staining images of positive cells of p-Src, p-VE-cadherin, p-Caveolin-1 (red staining), and CD31-positive cells (green staining) on paraffin-embedded sections of senescent C57BL/6 mice dorsal skin injected PBS (control) or MSC-sEV (scale bar, 200 μm). Quantitation of positive cells of p-Src, p-VE-cadherin, p-Caveolin-1 in CD31-positive cells. Representative immunofluorescence staining images of positive cells of p-Src, p-VE-cadherin, p-Caveolin-1 (red staining), and CD31-positive cells (green staining) on paraffin-embedded sections of type-2 diabetes model C57BL/6 mice dorsal skin injected PBS (control) or MSC-sEV (scale bar, 200 μm). Quantitation of positive cells of p-Src, p-VE-cadherin, p-Caveolin-1 in CD31-positive cells. Western blot analysis showing the phosphorylation change of Src, VE-cadherin, and Caveolin (downstream of Src) after transfected mimic-146a in HUVEC.
and also related to HG-induced endothelial inflammation control through IRAK-1 modulation. Moreover, miR-146a-5p partly contributed to the treatment of group 2 innate lymphoid cell-dominant allergic airway inflammation. Rau CS et al. discovered miR-146a may play a role in regulating the angiogenesis in HUVECs by downregulating CARD10, which acts in a negative feedback regulation loop to inhibit the activation of NF-kB that normally impairs angiogenesis. Su ZF et al. showed that miR-146a/b promoted proliferation, migration, and angiogenesis ability of endothelial progenitor cells through downregulation of TRAF6 and IRAK1 expressions. And Li Y et al. also found that miR-146a-mediated TGF inhibition, contributing to angiogenesis. However, most of these studies focused on the effect of cellular miR-146a. However, to our knowledge, no study has shown a link between MSC-sEV and miR-146a in the regulation of EC senescence. Altogether, our research suggests that miR-146a-5p may be a novel target for the treatment of cardiovascular diseases and diabetic vascular complications via rescue senescent ECs. We then investigated which protein influenced the senescent ECs through MSC-sEV. We found that among JNK1/2/3, Src, P38, AKT1/2/3, and Erk1/2 which we selected to increase the senescence of HUVECs, inhibition of Src decreased HUVECs senescence the most. The Src kinase family has been implicated in aging processes, Judith Haendeler et al. showed that the activity of the Src kinase family enhanced by ROS accumulation with age and induce lipid peroxidation, protein modification, DNA strand breaks, and oxidative damage, which results in a progressive loss of cell function, a hallmark for aging processes, and suggested that H2O2-induced ECs senescence was completely blocked with the Src-family kinase inhibitor PP2, which was consistent with our results. Sascha Jakob et al. also showed that Src could play a role in aging processes, since Src is responsible for TERT tyrosine phosphorylation, and Src deficiency can retain telomerase reverse transcriptase (TERT) in the nucleus under oxidative stress. We provided functional evidence that Src, the downstream of Src, VE-cadherin and Caveolin-1, were phosphorylated by senescence, and MSC exosomes inhibited the activation of Src both in H2O2 induced senescent HUVECs and high-glucose induced senescent HUVECs. This phenomenon had been identified in aged and type-2 diabetes C57BL/6 mice model. We also found that mimic-146a could inhibit the activation of Src kinase, which meant that MSC-sEV regulated the activation of Src partially through miR-146a. To find out how miR-146a regulate SRC, we first searched for the targets of miR-146a through miRTarBase, and we found 11 experimentally validated targets. However, Src is not among these validated targets, suggesting that miR-146a doesn’t regulate Src directly. We postulated that one or more molecules might be involved in the regulation between miR-146a and Src. Further experimental data are needed, and we will continue this investigation in the future. In short, our findings show that MSC-sEV can effectively increase senescent EC recovery and accelerate the healing of aged and diabetic mice wounds through increased angiogenesis. In the process of regulating functional recovery and wound healing dependent on MSC-sEV, miR-146a and Src play a crucial role, miR-146a can partially promote the pro-angiogenic and wound-healing effects, and subsequently dephosphorylate Src. Our findings suggest that by recovering senescent ECs via miR-146a, MSC-sEV may represent a promising strategy for vascular aging induced disease.

**MATERIALS AND METHODS**

**Cell culture**

Human adipose tissues and umbilical cords were obtained according to procedures approved by the Ethics Committee at Peking Union Medical College and the Chinese Academy of Medical Sciences. MSCs were isolated from human adipose tissues of healthy volunteers and culture-expanded as previously reported. HUVECs were isolated from human umbilical cords and cultured in endothelial cell medium (ECM #1001; ScienCell), following protocols described before. MSCs of passage 5 and HUVECs of passage 3 or 4 were used for our experiments. Cells were cultured in a humidified incubator with 5% CO2 at 37 °C and passaged with trypsin/EDTA after reaching the confluence.

**Transmission electron microscopy**

The morphology of MSC-sEV was examined by transmission electron microscope. On a carbon-coated copper grid, 20 µL of exosome suspension was simply loaded and set for at least 5 min. The MSC-sEV were subsequently stained with 2% uranyl acetate and dried for 10 min. Then, the grids were visualized with...
transmission electron microscope (Tecnai G2 Spirit TEM, Zeiss, Oberkochen, Germany) at 120 kV.

Nanoparticle tracking analysis
Nanoparticle Tracking Analysis (Zetaview, Particle Metrix) was used to evaluate the size distribution of MSC-sEV. Based on Brownian motion and diffusion coefficient, the particles are tracked and sized automatically. MSC-sEV is diluted to 1.0 mL with PBS. The measurement conditions for each sample were 23.75 ± 0.5 °C, 25 frames per second (FPS) for 60 s, with a similar detection threshold.

Exosomes internalization assay
MSC-sEV were labeled with a red fluorescent dye (PKH26; Sigma) according to the manufacturer’s instructions. The labeled sEV were then added to HUVECs or senescent HUVECs and co-cultured for 30 min, 2, 4, and 6 h. HUVECs were washed with PBS and fixed in 4 % paraformaldehyde for 15 min. Nuclei were stained with Hoechst, and the signals were analyzed with a fluorescence microscope.

To investigate mechanisms for internalization of MSC-sEV by HUVEC, we applied various endocytic inhibitors. Chlorpromazine (CPZ) is a widely used chemical inhibitor to interact with clathrin-mediated endocytic route (CME) by preventing clathrin lattices from being installed on endosomal membranes and clathrin-coated pits from being formed on the cell surface.

Nystatin is known to interfere with the caveolin-1 related uptake, which is associated with caveola formation.

We used CPZ (20 μM), nystatin (50 μM), and EIPA (50 μM) treated HUVECs for 30 min before the PKH26-labeled MSC-sEV were added.

Isolation of sEV from MSC
We followed the MISEV 2018 guidelines to isolate and identify MSC-sEV. Briefly, after MSCs reached 80–90% confluence, the serum-free medium was added for 48 h to avoid contamination of vesicles from serum. The conditioned medium was collected and centrifuged 800 g for 5 min and additional 3000 g for 10 min to remove cells and debris. The supernatant was then subjected to centrifugation to eliminate cell debris and large vesicles, followed by a 100,000-Mw cutoff membrane concentration (Centriplus-70, Millipore). The supernatant volume was reduced to less than 5 mL from approximately 250–500 mL. Using the 70Ti Rotor, the supernatant was then ultra-centrifuged at 110,000 g for 2 h at 4 °C (Beckman Coulter). The resulting pellets were resuspended with PBS and ultra-centrifuged with 100 Ti Rotor for 1 h at 110,000 g at 4 °C (Beckman Coulter). We used phosphate-buffered saline (PBS) as a negative control in the experiments involving MSC-sEV.

Induction of HUVECs senescence
H2O2 and high glucose were used to induce HUVEC senescence. Cells at 70% confluence were exposed to different doses of H2O2 (0, 25, 50, 75, and 100 μM) for 2 h and then washed with PBS and changed to fresh media. To get high-glucose-induced senescent HUVECs, we treated HUVECs with 30 mM d-glucose (HG) for 48 h. Control group was cultured in the media with normal glucose alone (5.5 mM).

Quantitative real-time polymerase chain reaction(qRT-PCR)
TRizol was used to extract the total RNA. At 260 and 280 nm, RNA concentrations and purity were calculated by optical density. Real-time PCR amplification was carried out in triplicate. SYBR Green RT-PCR (Takara Biotechnology Co., Ltd., Tokyo, Japan) was used to cDNA fragments for quantitative PCR. Relative mRNA expression was assessed using the 2-ΔΔCt method and normalized to GAPDH expression.

Western Blotting
Radioimmunoprecipitation (RIPA) lysis buffer with PMSF extracted proteins and the BCA Protein Assay Kit were used to quantify proteins. We performed Western Blot in triplicates followed the procedures reported previously and used GAPDH as the internal control.

Immunofluorescence staining
Cultured cells were fixed for 10 min at 4 °C in ice-cold methanol, washed three times in phosphate-buffered saline (PBS) and then at room temperature permeabilized in 0.1 percent Triton X-100/PBS for 10 min. With 0.5 percent Tween-20/PBS containing 1 percent bovine serum albumin (BSA) for 30 min, nonspecific binding was blocked. At 4 °C overnight, the primary antibodies were incubated. We used antibodies VWF (1:500; rabbit IgG) and CD31(1:500; rabbit IgG) for experiments. The secondary antibodies were, at room temperature, incubated for 1 h. PBS washed the incubated cells, and Hoechst was used to label the nuclei.

Transwell assay
We performed transwell assays by using 24-well transwell inserts with 8 μm pore-sized filters and 24-well culture plates (Corning, NY, USA). Cells were suspended in a low serum medium (containing 3% FBS) and plated into the upper chamber (1 × 10^4 cells per well; three replicates per group). In the lower chamber, 500 μL of complete medium (containing 10% FBS) were added. Cells attached to the upper surface of the filter membranes were removed using cotton swabs after 12 h of incubation and cells on the lower side of the filter (the migrated cells) were stained with crystal violet.

Scratch wound-healing assay
Cells (2 × 10^5 cells per well; three replicates per group) were seeded and incubated at 37 °C in a 12-well plate. We scratched the monolayer with a p200 pipette tip after the cells had been attached, washed with PBS to remove floating cells. At 0 h, 6 h and 12 h later, HUVECs were photographed. The migration area rate was determined as the closure area to initial wound ratio. Migration area (%) = (A1 – A0)/A0 × 100, where A1 represents the width of initial wound area and A0 represents the remaining width of wound at the metering point which analyzed by Image J software.

Tube formation assay in Matrigel
By conducting a tube formation assay, in vitro capillary network formation was determined in Matrigel. 1 x 10^5 HUVECs were plated in triplicates with 100 μL serum-free medium on a growth-factor-reduced Matrigel (BD)-coated 96-well plate. Tube formation was examined by microscopy after 6 h of incubation (Olympus, Tokyo, Japan) and branch points and tube length were quantified by choosing three fields per well at random with Image J software.

Enzyme linked immunosorbert assay (ELISA)
HUVECs conditioned medium was collected from each group (n = 3 per group) after PBS or MSC-sEV co-cultured for 24 h, stored at room temperature for 15 min and then centrifugated at 2000 g for 10 min. The samples were either frozen at −80 °C or immediately analyzed using human ELISA kits for IL-1alpha, IL-6, and IL-8. We performed all procedures by following the manufacturer’s instructions and measured the absorbance at 450 nm by using a microplate reader (Bio-Rad, USA). Each determination was the average of at least three independent experiments.

Mitochondrial respiration analysis using the Seahorse XF Analyzer
To calculate the oxygen consumption rate (OCR), the Mito Stress Test Kit was used. The probe plate was hydrated with HPLC grade water in a CO2-free incubator prior to metabolism calculation.
In order to preserve the pH value, the test phenol red-free solution containing 10 mM glucose, 2 mM glutamine, 1 mM pyruvate and 5 mM HEPES was kept in a 37 °C CO₂-free incubator. In the hydration plate, the HPLC grade water was then replaced with a calibration solution and stored in a CO₂-free 37 °C incubator. When density of cells was at 5000 per well, HUVECs were seeded into a 96-well plate at the density of 3 × 10⁵ cells per well. After 24 h, cells were washed with PBS three times to remove unbound dye and photographed under a light microscope.

Reactive oxygen species (ROS) detection

Intracellular ROS generation in HUVECs was measured by using Fluorometric Intracellular ROS Kit. The HUVECs of each group (n = 3 per group) were seeded onto 6-well plates at the density of 3 × 10⁵ cells per well in the culture medium. After the cells reached 80% confluence, they were subjected to H₂O₂ stimulation, and then incubated with MSC exosomes for 12 h. After the indicated time, the cells were incubated with ROS Kit for 30 min at 37 °C. Finally, these cells were washed with pre-warmed PBS and tested using a flow cytometer (BD, Accuri C6 Cytometer, USA) within 1 h. We performed each experiment at least for 3 samples and each determination was made in triplicate.

Proliferation assay

We seeded cells into 96-well culture plates for 5 × 10³ cells per well and each group had four replicates. A no-cell community calibration solution and stored in a CO₂-free 37 °C incubator. In the hydration plate, the HPLC grade water was then replaced with a red-free phenol assay solution and put for 1 h in a 37 °C CO₂-free incubator. We measured the absorbance minus the blank value of each well. We performed each experiment at least for 3 samples and each determination was made in triplicate.

Mouse skin wound model and treatments

We purchased aged female C57BL/6 J mice normal male C57BL/6 J from the Laboratory Animal Center of Peking Union Medical School (Beijing, China). The mice were 64-week old when we separated them into two groups at random (n = 6): PBS group (treated with 100 μL PBS), MSC-sEV group (treated with 100 μL PBS). Briefly, the mice were subcutaneously injected with MSC-sEV or PBS around the wounds at 4 injection sites (25 μL per site). When on days of 0, 3, 6, 9, and 12 post-wounding, wounds were photographed and measured. At 12 days post operation, mice were sacrificed, and skin samples were harvested and fixed in 10% paraformaldehyde (PFA). Samples from each group were subjected to hematoxylin eosin (H&E) staining for the detection of newly formed blood vessels.

Matrigel plug assay

Growth-factor-reduced Matrigel aliquots (150 μL) containing 5 × 10⁶ HUVECs pretreated with H₂O₂ or MSC-sEV were prepared on ice. Mice were randomly divided into three classes (control, 5 and MSC-sEV, n = 3) and anaesthetized with a 50 mg/kg pentobarbital sodium intraperitoneal injection (Sigma Aldrich). Matrigel aliquots were injected into the inguinal areas bilaterally and could gel at body temperature. The mice were euthanized for plug excision after 14 days. Plugs were fixed with 4 % PFA and stained with anti-human CD31 for angiogenesis detection.

Transfection of mimic and inhibitor

HUVECs in 6-well plates (about 5 × 10⁵ cells/well) were transfected with miR-146 mimic (50 nmol/L), or miR-146 inhibitor, or miR-34 inhibitor (100 nmol/L), or their corresponding negative controls. Lipo2000 transfection reagent was simultaneously added into the medium for efficient transfection. After 6 h, we replaced the culture medium in order to remove the transfection reagent. Detection was made 24 h after transfection. miR-146a mimic, miR-146a inhibitor, miR-34a inhibitor, and their negative controls were purchased from Sangon Biotech.

Oil red O staining, Alizarin red staining and Alcian Blue staining

After lineage differentiation, cells were washed twice with PBS and fixed with 4 % formalin for 10 min at room temperature. For Oil red O staining, fixed cells were then stained with Oil red O reagent (Beyotime) for 1 h at room temperature; for Alizarin red staining, fixed cells were stained with 1% Alizarin red (Leagene, Beijing, China) with pH 4.2 for 30 min at room temperature; for Alcian Blue staining, fixed cells were then colored for 30 min in a 1% Alcian Blue solution (OrCell). Finally, cells were washed with water for three times to remove unbound dye and photographed under light microscopy.

Flow cytometry analysis

MSC immunophenotype analysis was performed as previously reported. Briefly, MSCs were washed and incubated with primary antibodies (CD29, CD34, CD44, CD45, CD73, CD90, CD105, CD106, CD206, and HLA-DR, BD Biosciences) for 30 min at 4 °C. After washing, secondary antibodies were used to incubate with MSCs for 30 min at 4 °C, and MSCs were analyzed by flow cytometer and CFlow Plus software.

Statistical analysis

We presented mean ± SD for all data. We analyzed the comparisons between groups using Student’s t test, one-way ANOVA followed by Tukey’s multiple comparisons test and Dunnett’s multiple comparisons test. Statistically significant differences were considered at *p < 0.05, **p < 0.01, ***p < 0.001.

DATA AVAILABILITY

All data supporting this paper are present within the paper and/or the Supplementary Materials. The original datasets are also available from the corresponding author upon request.

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AUTHOR CONTRIBUTIONS
XX. performed and analyzed experiments, prepared the figures, and wrote the manuscript. W.S. designed and analyzed experiments and wrote the manuscript. X.M., Y.H., W.L., & L.X. performed the experiments. J.R. and Z.C. designed the experiment. All authors have read and approved the final manuscript.

ADDITIONAL INFORMATION
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