A pH probe inhibits senescence in mesenchymal stem cells

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Abstract

Background: Bone marrow-derived mesenchymal stem cells (BMSCs) are gradually getting attention because of their multi-directional differentiation potential, hematopoietic support, and promotion of stem cell implantation. However, cultured BMSCs in vitro possess a very limited proliferation potential, and the presence of stem cell aging has substantially restricted the effect together with the efficiency in clinical treatment. Recently, increasing attention has been paid to the connection between cellular aging and lysosomal acidification as new reports indicated that vacuolar H⁺-ATPase (v-ATPase) activity was altered and lysosomal pH was dysregulated in the process of cellular aging. Therefore, promoting lysosomal acidification might contribute to inhibition of cell senescence. Our previous studies showed that a novel small molecule, 3-butyl-1-chloro imidazo [1, 5-a] pyridine-7-carboxylic acid (SGJ), could selectively and sensitively respond to acidic pH with fast response (within 3 min), but whether SGJ can promote lysosomal acidification and inhibit senescence in BMSCs is unknown.

Methods: Rat BMSCs were cultured based on our system that had been already documented. BMSCs were treated with SGJ and/or Bafilomycin-A1 (Baf-A1). The co-localization between SGJ and lysosomes was assessed by a confocal microscope. Acridine orange (AO) staining and the Lysosensor™ Green DND-189 reagents were used for indicating changes in lysosomal concentration of H⁺. Changes of senescence were detected by immunoblotting of p21 and senescence-associated beta-galactosidase (SA-β-gal) staining as well as immunofluorescence assay of senescence-associated heterochromatin foci (SAHF). Changes of autophagy were detected by immunoblotting of MAP1LC3 (LC3B) and SQSTM1 (p62). Cell proliferation was determined by flow cytometry. Cell viability was calculated by sulforhodamine B assay (SRB). The V0 proton channel of v-ATPase was knocked down by transfecting with its small interfering RNA (si-ATP6V0C).

Results: Our work showed that SGJ can promote lysosomal acidification and inhibit senescence in BMSCs. Firstly, SGJ and lysosomes were well co-located in senescent BMSCs with the co-localization coefficient of 0.94. Secondly, SGJ increased the concentration of H⁺ and the protein expression of lysosome-associated membrane protein 1 (LAMP1) and lysosome-associated membrane protein 2 (LAMP2). Thirdly, SGJ suppressed the expression of p21 in the senescent BMSCs and reduced SA-β-gal positive cells. Fourthly, SGJ promoted senescent BMSCs’ proliferation and protein level of LC3B but reduced the p62/SQSTM1 protein level. Furthermore, experimental group pretreated with 20 μM SGJ showed a stronger red fluorescent intensity, thinner cell morphology, less SA-β-gal positive cell, and less p21 protein level as well as higher cell viability in the presence of Baf-A1. Notably, ATP6V0C knockdown decreased the activity of v-ATPase and SGJ increased the concentration of H⁺.

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Background

An important member of stem cell family, bone marrow-derived mesenchymal stem cell (BMSC) that can be induced to differentiate into adipocytes, osteocytes, muscle cells, nerve cells and endothelial cells, originated from the early mesoderm, are gradually getting attention because of its multi-directional differentiation potential, hematopoietic support and promotion of stem cell implantation [1–3]. The application of BMSCs in clinical infusión is now providing a great hope for cell therapies and tissue engineering in a wide variety of diseases [4–7]. We think that BMSCs can be used as an ideal seed cell for aging and lesions in the future [8, 9]. However, cultured BMSCs in vitro possess a very limited proliferation potential, and the presence of stem cell aging has substantially restricted the effect together with the efficiency in clinical treatment, especially impeding the development of an autologous cell-based therapy for senior patients [10].

Cellular senescence involves an extensive range of blocking cellular remodeling and cycling, including chromatin structure [11–13]. It has been suggested that the increased intracellular balance failure during cell aging was partly due to the impaired function of lysosome that was the key organelle for autophagy, and the presence of lysosome and autophagy was important to cellular constituents recycling and cell remodeling involved in cellular senescence [1, 14–18]. In addition, metabolites and ions released by lysosomes play a vital role in signal transduction and nutritional sensation, which are affected by intraluminal pH of lysosomes [19, 20]. Lysosomes maintain an acidic lumen by means of the proton pump and the vacuolar H+-ATPase (v-ATPase) [21]. H+-ATPases are large, multi-subunit complexes composed of a peripheral domain (V1) that hydrolyzes ATP and a membrane integral domain (V0) that translocates protons into lysosomes and therefore lowers the intraluminal pH to the acidic range [22]. With the acidic pH optimum conditions, dozens of hydrolytic enzymes in the lysosome are activated [23]. Therefore, the disorders of lysosomal acidification can have obvious effects on lysosomal digestion. The disorders strongly inhibit the activity of acidic hydrolases, but they also potentially promote the activities of other neutral hydrolases [24]. This transformation will promote both substrate indigestion and atypical cleavages, which may also produce toxic products and/or partially digested intermediates [1]. Increasing attention has been paid to the connection between cellular aging and lysosomal acidification as new reports are indicating that v-ATPase activity is altered and lysosomal pH is dysregulated in the process of cellular aging [1]. The disorders of lysosomal acidification increase the risk of cellular aging and death [1]. However, vacuolar acidification appears to be critical in mediating the prolonged lifespan effects of caloric restriction and methionine restriction [25–27]. In addition, overexpressing of the vacuolar ATPase (v-ATPase) components promotes an increase in lifespan in the yeast models [25, 27]. Increased vacuolar ATPase (v-ATPase) early in life contributes to age-related mitochondrial dysfunction. It is reasonable to expect that upregulated function of v-ATPase that promotes lysosomal acidification contributes to inhibition of cell senescence [25–27]. Thus, successful applications of small chemical molecules in inhibiting senescence in BMSCs by maintaining lysosomal functions provide potential intervention strategies to prevent body from aging and regenerate tissues [28]. Our previous work showed that a novel small molecule, 3-butyl-1-chloro imidazo [1, 5-a] pyridine-7-carboxylic acid (SGJ), had been demonstrated as a fluorescent probe to selectively and sensitively respond to acidic pH with fast response (within 3 min) [29]. Here, we wish to report our studies on SGJ, in which clarify its functions in lysosomes, relationship with Baf-A1 and role in suppressing cell senescence.

Methods

Cell extraction and culture

Rat bone marrow collection and mesenchymal stem cell isolation were performed according to Pittenger et al. In brief, bone marrow in the femurs and tibias of male Wistar rats (90–100 g) was washed by using Dulbecco’s modified Eagle’s medium-low glucose (DMEM-LG) (Gibco) supplemented with 10% fetal bovine serum (HyClone). BMSCs were isolated by removal of non-adherent cells after 72 h, and the medium was changed every 2 days until the adherent cells reached 50% confluence, and they were then harvested with 0.05% trypsin (Sangon Biotech) in phosphate-buffered saline (PBS). Cells were grown to 70–80% confluence and then seeded on plates or appropriate dishes at the density of 5000 cm$^{-2}$. The experiments were conducted on cells at varied

Conclusion: Our work showed that SGJ could inhibit senescence in BMSCs and protect lysosomes by promoting expression of LAMP1 and LAMP2. Meanwhile, SGJ could promote autophagy. Furthermore, our study also suggested that SGJ was a new Baf-A1 antagonist because SGJ could target and occupy the V0 proton channel of v-ATPase.

Keywords: Autophagy, Bafilomycin-A1 antagonist, Senescence, Bone marrow-derived mesenchymal stem cells
population doubling level (PDL) to demarcate young (PDL 3–6) and senescent (PDL 16–25) BMSCs.

**Cell treatment**

BMSCs were cultured in Dulbecco’s modified Eagle’s medium-low glucose (DMEM-LG) (Gibco) supplemented with 10% fetal bovine serum (HyClone) at 37 °C in a humidified incubator with 5% CO₂. In the process of the whole experiments, the cell’s density was around 90% before doing all experiments, and the decentralized cells were seeded onto 6-well plates, 12-well plates, 24-well plates, 96-well plates, or other appropriate dishes at the density of 20,000 cells/mL with pre-incubation for 12 h before adding the compound SGJ. SGJ was synthesized as reported with a purity greater than 98% and the structure was confirmed by spectral data. SGJ was dissolved in dimethylsulfoxide (DMSO) to make a 0.1 M stock solution. BMSCs were incubated with culture medium containing 1, 5, 10, and 20 μM SGJ, or a corresponding volume of DMSO solvent as control (below 0.1%, v/v) in all subsequent experiments.

**Cell morphological observation**

BMSCs were seeded onto 24-well plates and treated with 0.1% DMSO as a negative control or 1, 5, 10, and 20 μM SGJ for 24 or 48 h, respectively, and the microscopic photographs (×200) were taken under the inverted phase-contrast microscope (Nikon, Japan).

**Cell viability assay (SRB)**

Referring to the relevant reports, BMSCs were seeded onto 96-well plates. After 12 h, cells were respectively treated with 0.1% DMSO as a relevant control or compound SGJ at 1, 5, 10, and 20 μM for specified time durations. Then the cell viability was evaluated by sulforhodamine B assay (SRB), according to the method of Skehan. In short, discarded the original medium and added 100 μL cold 10% trichloroacetic acid (TCA) into the 96-well plates to fix cells and incubated for 1 h at 4 °C. Poured off the supernatant and washed the cell five times with ultrapure water. After drying the 96-well plates at room temperature, we added 50 μL of 0.4% (w/v) SRB solution in 1% acetic acid into each hole and shook it for 10 min on a microtiter plate shaker. Then we washed the 96-well plates five times with 1% acetic acid and subsequently added 100 μL of 10 mM Unbuffered Tris Base (pH = 10.5) to dissolve the original dye after again drying the 96-well plates at room temperature. We shook the plate on a microtiter plate shaker for 10 min and measured the light absorption at the wavelength of 540 nm by a SpectraMAX190 microplate spectrophotometer (GMI Co, USA).

**Flow cytometry**

Cells were digested into single cells by 0.25% trypsin (Sangon Biotech) and collected into 15-mL centrifuge tubes. Centrifugation at 1000 rpm for 5 min was performed, then we discarded the supernatant. We washed cells twice with 1× PBS buffer and suspended them in 1× PBS buffer by centrifugation at 1000 rpm for 5 min each time. We discarded the supernatant. We suspended cells by 1 mL cold 70% ethyl alcohol and fixed them at 4 °C overnight. We washed cells three times with 1× PBS buffer by centrifugation at 1000 rpm for 5 min each time. We discarded the supernatant and suspended 1×10⁶ or 1×10⁷ cells with 500 μL 1× PBS buffer including 100 μg/mL propidium iodide (PI) (Biolegend), 50 μg/mL RNAase (TIANGEN), and 0.2% X-Triton 100 (Sangon Biotech), and placed it at 4 °C for 30 min. It was analyzed by flow cytometry.

**Senescence-associated β-galactosidase assay**

BMSCs seeded in 24-well plates were exposed to 2% formaldehyde/0.5% glutaraldehyde for fixation. After 5 min, cells were rinsed with PBS and warmed β-gal buffer solution (5 mM potassium ferrocyanide, 5 mM potassium ferricyanide, 150 mM NaCl, and 2 mM MgCl₂, pH = 6.0) for 1 min. Then, cells were incubated with staining solution (buffer solution supplemented with 1 mg mL⁻¹ 5-bromo-4-chloro-3-indoly β-D-galactopyranoside, Takara) for over 18 h at 37 °C. Senescent cells were stained to be blue under a phase-contrast microscope. The percentage of positively stained cells was estimated by counting at least 1500 cells in each sample.

**Lysosomal activity detection**

Lysosomal activity was monitored by a metachromatic fluorophore acridine orange (AO) (0.1 mg mL⁻¹). After treatment, BMSCs were gently rinsed twice with PBS and then subjected to acridine orange staining for 1 min. Cells were washed with PBS and the fluorescence was observed under an inverted fluorescence microscope (Nikon).

**Immunofluorescence microscopy**

For the next immunofluorescence assay, we seeded BMSCs onto confocal dishes (20 mm) (SPL, Korea) and used SGJ to treat cells. After treatment, we aspirated the culture medium and rinsed the cells gently with 1× PBS, then the cells were fixed in 4% paraformaldehyde for 15 min. After washing three times with 1× PBS, permeated cells with 0.1–0.2% TritonX-100 for 5 min, then washed and blocked with donkey serum (1:30 dilution in 0.1 M PBS) for 20 min at room temperature. We discarded the enclosed liquid, incubated cells with primary HP1-γ antibody (Proteintech), H3K9Me3/2 antibody (Cell signaling, USA) (1:100 dilution) overnight at 4 °C, then incubated cells with Alexa 488 nm labeled species-specific secondary
antibodies (1:200 dilution in 0.1 M PBS) for 60 min at 37 °C. Finally, we rinsed and stained cells with DAPI and monitored them by a confocal laser scanning microscope (Zeiss, Germany).

GFP-LC3B puncta observation
U87 cells were seeded onto confocal dishes (20 mm) (SPL, Korea) and used SGJ and chloroquine (CQ) to treat cells. After treatment, we discarded the culture medium and rinsed the cells gently with 1× PBS, then the cells were fixed in 4% paraformaldehyde for 15 min. After washing three times with 1× PBS, we observed the cells with a confocal laser scanning microscope (Zeiss, Germany).

Western blot analysis
As described previously, cells were treated with 0.1% DMSO or SGJ for specific time and washed twice with ice-cold PBS, then lysed in protein lysis buffer (Shanghai Beyotime Co, China) that contained 0.5% SDS in 25 mM Tris–HCl, 4 mM EDTA, 100 mM NaCl, 1 mM PMSF, 10 μg mL⁻¹ leupeptin, and 10 μg mL⁻¹ soybean trypsin inhibitor, pH = 7.5. The protein concentrations were determined by the Bradford method. Before being loaded into a 12% or 15% SDS polyacrylamide gel, equal amounts of proteins were added with loading buffer and boiled. Following electrophoresis, the resolved protein was electrophotore transfered to a polyvinylidene fluoride (PVDF) membrane (Millipore, MA, USA). The membrane was blocked with 5% non-fat milk in TBST (Tris buffer saline containing 0.5% Tween 20) for 1 h at room temperature. Subsequently, the membrane was probed with p21 antibody (1:2000) (Proteintech, USA), LC-3B antibody (1:2000) (Cell signaling, USA), p62 antibody (1:2000) (BOSTER, USA), LAMP1 antibody (1:500) (SIGMA, USA), or anti-β-actin mouse monoclonal antibody (1:2000) (Santa, USA) overnight at 4 °C, then it was washed with TBST for 5 min twice. The membrane was subsequently incubated with HRP-conjugated goat anti-rabbit IgG (1:5000) (Beijing Dingguo Co, China) or polyclonal goat anti-mouse immunoglobulins/HRP (1:5000) (Beijing Dingguo Co, China) for 1 h at room temperature and then washed three times with TBST. Then the membrane was incubated with HRP substrate for 5 min, and the fluorescence signal was detected with X-ray films. Intensity of the protein bands was quantified by Quantity-One software (Bio-Rad), analyzed by ImageJ software, and normalized to loading controls.

Lysosomal co-localization experiment
We seeded BMSCs onto confocal dishes (SPL, Korea). After 12 h, cells were treated with 0.1% DMSO (as control) or compound SGJ (20 μM) for 1 h, and then they were treated with 0.5 μM LysoTracker® Red DND-99 for 30 min. We washed cells with 1× PBS for three times and monitored them by a confocal laser scanning microscope (Carl Zeiss, Germany). The excitation wavelength of SGJ is 380 nm, while the excitation wavelength of LysoTracker® Red DND-99 is 577 nm.

Lysosomal pH sensing experiment
We seeded BMSCs onto confocal dishes (SPL, Korea). After 12 h, the cells were treated with 0.1% DMSO (as control), compound SGJ (20 μM), and/or Bafilomycin-A1 (Baf-A1) (20 nM) for a specific time, and then cells were treated with 0.5 μM Lysosensor™ Green DND-189 (pH = 5.2) for 30 min. We washed cells with 1× PBS three times and monitored them by a confocal laser scanning microscope (Carl Zeiss, Germany). The excitation wavelength of Lysosensor™ Green DND-189 is 443 nm.

Si-RNA interference experiment
The specific siRNA against ATP6V0C was synthesized by Invitrogen. Scramble siRNA was used as a control (Santa Cruz, sc-37007). The corresponding negative control (NC) was designed and purchased from Invitrogen. Cells at 70% confluence were transfected with siRNA by Lipofectamine 2000 (Invitrogen, 11668–019) transfection reagent according to the manufacturer’s instructions.

Statistical analyses
Data were presented as means ± SE from no less than three independent experiments and analyzed by SPSS (Statistical Package for the Social Sciences) software and Student’s t test. Pictures were processed with Adobe Photoshop software. The mean values were derived from at least three independent experiments. Differences at p < 0.05 were considered statistically significant.

Results
SGJ co-located with lysosomes
The chemical structure of the small molecule SGJ is shown in Fig. 1a. To explore the relation between SGJ and lysosome, we treated BMSCs with SGJ and LysoTracker® Red DND-99. We found that SGJ and lysosome are well co-located in senescent BMSCs with the co-localization coefficient of 0.94 (the complete co-localization coefficient is 1) (Fig. 1b).

SGJ increased the concentration of H⁺ and protected the function of lysosomes in senescent BMSCs
Wang et al. proved that lysosomal activity declined and acidic vacuoles decreased with age [28]. Acridine orange (AO) is normally used as an indicator for changes in lysosomal pH, lysosomal integrity and permeability [30, 31]. As shown in Fig. 2a, to clarify the function of SGJ in lysosome, we performed AO staining. The results showed that the senescent cells at PDL 20 displayed a
dim red fluorescence compared to the young cells at PDL 5. SGJ treatment greatly increased red puncta in the senescent cells, indicating a higher level of acidic vacuoles. To investigate whether SGJ functioned by increasing the concentration of H⁺ in the senescent lysosome or not, we next utilized Baf-A1, a recognized inhibitor of v-ATPase located on the lysosomal membrane, which reduces the concentration of H⁺ in lysosome. We exposed the senescent cells to 20 nM Baf-A1 and 20 μM SGJ for 24 h, then, incubated the cells with Lysosensor™ Green DND-189 and photographed under the confocal microscopy. The Lysosensor™ Green DND-189 reagents exhibit a pH-dependent increase in fluorescence intensity upon acidification. We found that the red fluorescence nearly disappeared in the control cells, while it further decreased in the Baf-A1 treated cells and obviously increased in the SGJ treated cells (Fig. 2b). The results indicated that SGJ could increase the concentration of H⁺ in lysosome in the senescent BMSCs.

Immunoblotting of LAMP1 (lysosome-associated membrane protein 1) and LAMP2 (lysosome-associated membrane protein 2), the established biomarkers of the function of lysosome, showed a decreased level of these markers in senescent cells. Treating senescent BMSCs with SGJ significantly increased the protein expression of LAMP1 and LAMP2, which restored the lysosomal activity (Fig. 2c). Our results indicated that SGJ might have a protective function on damaged lysosomes.

**SGJ inhibited cultured BMSC senescence and promoted BMSC proliferation**

We then discovered that SGJ treatment significantly suppressed the senescent biomarkers in BMSCs (Fig. 3). The SA-β-gal staining result showed that there were more positively stained cells in the senescent group (PDL 22) than in the young group (PDL 6) (Fig. 3a). Meanwhile, we found an obvious morphological change in senescent BMSCs. Comparing with the young cells’ spindle-type morphology, the morphology of the senescent BMSCs showed a more flat and spread out status, and the phenomenon of podia and actin stress filaments was increased [32]. We found that using SGJ to treat the senescent BMSCs can significantly decrease the number of SA-β-gal positive cells compared to the control group (PDL 22) (Fig. 3a). SGJ-treated senescent BMSCs exhibited a thinner and smaller morphology. The cytoplasmic extensions and the actin filaments were decreased. The cytoplasmic extensions and the actin filaments were decreased. Senescence-associated heterochromatin foci (SAHF) are specialized domains of facultative heterochromatin related to the contribution to silencing of proliferation-promoting genes in senescent cells. SAHF and its components such as heterochromatin protein 1 (HP1) proteins and lysine 9 di-or tri-methylated histone H3 (H3K9Me2/3) are always used as markers for senescent cells [33]. We found that, compared to the young cells (PDL 6) there were robust punctate DAPI foci in the senescent BMSCs (PDL 22), which co-localize with both HP1 gamma homolog (HP1-γ) and lysine 9 di-or tri-methylated histone H3 (H3K9Me2/3). However, treatment with SGJ for 48 h decreased the punctate DAPI foci and its co-localization with both HP1-γ and H3K9Me2/3 in senescent BMSCs (PDL 22) (Fig. 3b). We also found that SGJ treatment significantly decreased the protein level of p21, an established marker of aging in senescent BMSCs (Fig. 3c).
Next, to find out effects of SGJ on the senescent BMSCs’ growth kinetics, cells (PDL22) were treated with SGJ for 48 h and then stained with propidium iodide (PI) and analyzed by flow cytometry (Fig. 3d). We found about 80.14% senescent BMSCs (PDL 22) stayed at G0/G1 phase and about 19.86% stayed at S phase or G2/M phase (Fig. 3d), while in SGJ-treatment group, the percentage of cells staying at G0/G1 phase decreased to about 68.85% and cells at S phase or G2/M phase increased to 31.15% (Fig. 3d). Also, we found that SGJ could significantly increase the viability of senescent BMSCs (Fig. 3e). Besides, the density of BMSCs increased significantly with the morphology of BMSCs becoming thinner and plumper than control group (PDL 22) (Additional file 1: Figure S1). Taking all these results into account, we can draw a conclusion that SGJ suppresses BMSCs aging in vitro, and promotes the senescent BMSC proliferation. Meanwhile, autophagy plays an important role in the mechanisms of cellular aging. Recently, it is exemplified by studies revealing that increased autophagy flux is a potential mechanism for extending life-span in yeast, drosophila and mouse models. Thus, we investigated the effects of SGJ on BMSC autophagy. We found that SGJ increased the level of protein LC3B significantly and reduced the p62/SQSTM1 protein level by promoting autophagic degradation. To validate the effect of SGJ on autophagy flux, we used Baf-A1/chloroquine (CQ) to block the flow of autophagy (Additional file 1: Figure S2). The data showed that SGJ promoted the flow of autophagy in
BMSCs (Additional file 1: Figure S2). It also revealed that treating BMSCs with SGJ could induce cell autophagy significantly.

**SGJ protected and enhanced lysosomal activity via protecting the V0 proton channel of v-ATPase**

The vacuolar ATPase (v-ATPase) is a macromolecular complex. It is responsible for pumping protons (H⁺) into lysosomes and lowering intraluminal pH to the acidic range, and it is needed to activate dozens of hydrolases with acidic pH optima in lysosomes [23]. We have already discovered that SGJ raised the concentration of H⁺ in lysosome in the senescent BMSCs. We wondered if the SGJ-increased concentration of H⁺ in lysosome was a consequence of protecting v-ATPase. The Baf-A1 inhibition site localizes to the V0 proton channel and requires residues of subunit C [34–36]. We next utilized Baf-A1 and / or SGJ to explore the targeting sites of SGJ. We designed three typical experiments: (A) treatment with 20 nM Baf-A1 for 6 h and then treatment with 20 μM SGJ for 18 h; (B) treatment with 20 μM SGJ for 6 h and then treatment with 20 nM Baf-A1 for 18 h; (C) co-treatment with 20 nM Baf-A1 and 20 μM SGJ to the senescent BMSCs for 24 h. Next, we detected the red fluorescence intensity by Lysosensor™ Green DND-189 staining (Fig. 4a), imaged the cells’ morphologic changes and calculated the cell viability by SRB (sulforhodamine B assay) (Additional file 1: Figure S3). We found that the experimental group pretreating with 20 μM SGJ showed a stronger red fluorescent intensity, thinner cells’ morphology and higher cell viability in the presence of Baf-A1. The results indicated that SGJ might occupy the action sites of Baf-A1 competitively, which targeted the V0 proton channel of v-ATPase to protect its activity, resulting in increasing the concentration of H⁺ and preventing the destructive effect of Baf-A1 in lysosome. For further exploring the possible molecular mechanism of SGJ in increasing the concentration of H⁺, 293 T cells were transfected with si-ATP6V0C at 20 or 60 nM for
24 h. Si-ATP6V0C could markedly decrease the activity of the V0 proton channel of v-ATPase. The efficiency of ATP6V0C knockdown was verified (Fig. 4b). The fluorescent intensity results showed that si-ATP6V0C decreased the activity of v-ATPase and SGJ increased the concentration of H⁺ (Fig. 4c).

Next, to further prove the targeting sites of SGJ is the V0 proton channel of v-ATPase. We added SGJ and/or Baf-A1 in a different order and detected the changes in SA-β-gal positive cells as well as the p21 protein level. Compared with other experimental groups, we found that the experimental group pretreating with 20 μM SGJ showed a fewer SA-β-gal positive cells and lower p21 protein level in the presence of Baf-A1 (Fig. 5). The data revealed that SGJ is able to occupy the action site of Baf-A1 preferentially and prevent cell senescence effectively.

**Discussion**

In the process of cell aging, the lysosomal activity is decreased and the concentration of H⁺ in lysosome is reduced [1, 28, 37]. Meanwhile, the autophagic ability of the senescent cells is reduced [13, 38, 39]. In this study, we discovered that the novel small molecule, 3-butyl-1-chloro imidazo [1, 5-a] pyridine-7-carboxylic acid (SGJ) could co-locate with lysosome and protect its function. We also found that SGJ could bind to v-ATPase, promote the inflow of H⁺ in lysosomes and inhibit the decrease in the concentration of H⁺ in senescent BMSCs. LAMP1 and LAMP2 are functional proteins on the lysosomal membrane, and they have been linked to protection of membrane from degradation by lysosomal hydrolases [40]. However, both the lysosomal activity and the protein levels of LAMP1 and LAMP2 decreased in the process of cell aging [40, 41]. Protein levels of LAMP1 and LAMP2 increased after treatment with SGJ, showing that SGJ protected the lysosomal integrity and function. The cellular lysosomes maintain a low acidic lumen by means of the proton pump and v-ATPase [21]. With the acidic pH optimum conditions, dozens of hydrolytic enzymes in the lysosome are activated [23]. However, new reports
implicate lysosomal pH dysregulation in cellular aging [1].
Therefore, an abnormal rise in lysosomal pH can have sig-
nificant effects on lysosomal digestion, for example,
strongly inhibiting hydrolases with the most acidic pH op-
tima, or potentially elevating other hydrolases’ activities
with pH optima being closer to neutral [23]. SGJ can pro-
mote the function of lysosomes themselves by elevating
activities of hydrolases with the most acidic pH optima.
Thus, this indicates that the functions of lysosomes re-
stored and the levels of cell autophagy are elevated in the
senescent cells under treatment with SGJ.

In fact, to promote the efficiency of BMSCs for clinical
therapies and tissue engineering, we should pay attention
to both intrinsic aging of BMSCs in vivo and their aging
in vitro when these cells may require expansion in vitro
before use as well as for future research [42]. It is no
doubt that there exist differences between PDL-based
aging model in vitro used in our research and those natur-
ally senescent BMSCs in vivo. For instance, in vivo, stem
cell fate and activities are largely determined by their
microenvironment. However, when cultured in vitro with-
out the presence of regulatory cues which exist in vivo,
MSCs may have different functions and properties [43].
When viewed from this perspective, whether SGJ could
exert the same effect on those BMSCs aged in vivo should
be investigated further. However, many studies indicated
that elements changed during aging, including autophagy,
lysosomal function and activity, and v-ATPase activity,
were not only found in vivo [1, 15, 38, 44] but also in vitro
[14, 15, 44–46]; therefore, according to our work, it is rea-
sonable to draw a conclusion that SGJ is a potential lead
compound for suppressing BMSC senescence. At least, it
could be used as a powerful tool for studying and under-
standing the mechanism of aging.

**Conclusions**
Our work showed that through treating the senescent
BMSCs with SGJ, the condition of cell aging is inhibited
and the viability of the senescent cells is improved. SGJ is a potential lead compound for suppressing BMSCs senescence. In addition, we further explored the targeted site of SGJ. By adding SGJ and/or Baf-A1 in a different order, we found that SGJ could target and occupy the V0 proton channel of v-ATPase, indicating that SGJ and Baf-A1 have a competitive relationship and SGJ protects the v-ATPase activity via preventing the destructive effect of Baf-A1 on lysosome (Fig. 4c). The results suggested that SGJ was a new Baf-A1 antagonist that promoted autophagy and inhibited senescence in bone marrow-derived mesenchymal stem cells (Fig. 6).

Additional file

Additional file 1: Supplementary figures are included in it. (DOC 1364 kb)

Abbreviations
AO: Acridine orange; Baf-A1: Bafilomycin-A1; BMSCs: Bone marrow-derived mesenchymal stem cells; CQ: Chloroquine; DMEM-LG: Dulbecco’s modified Eagle’s medium-low glucose; DMSO: Dimethylsulfoxide; H3K9Me2/3: Lysine 9 di-or tri-methylated histone H3; HP1: Heterochromatin protein 1; HP1-γ: HP1 gamma homolog; LAMP1: Lysosome-associated membrane protein 1; LAMP2: Lysosome-associated membrane protein 2; LC3B: MAP1LC3; p62: SQSTM1; PBS: Phosphate-buffered saline; PDL: Population doubling level; PI: Propidium iodide; PVDF: Polyvinylidene fluoride; SAHF: Senescence-associated heterochromatin foci; SA-β-gal: Senescence-associated beta-galactosidase staining; SGJ: 3-Butyl-1-chloro imidazo [1,5-a]pyridine-7-carboxylic acid; si-ATP6V0C: Small interfering RNA; SRB: Sulforhodamine B assay; TCA: Trichloroacetic acid; v-ATPase: Vacuolar H+-ATPase

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Availability of data and material
All data generated or analyzed during this study are included in this published article [and its supplementary information files].

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Authors’ contributions
JYM and BXZ designed the experiments. LHW, XJH, and GJQ conducted the experiments. LHW, XJH, and LS acquired the data. LHW, XJH, JYM, and BXZ analyzed the data. LHW, JYM, and BXZ organized the manuscript. All authors read and approved the final manuscript.

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Not applicable.

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Competing interests
The authors declare that they have no competing interests.

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Fig. 6 Schematic presentation of SGJ preventing the disorders of lysosomal acidification via v-ATPase in senescent BMSCs. In the process of cell aging, the lysosomal activity is decreasing and the concentration of H + in lysosome is reducing. SGJ can bind to v-ATPase, promote the inflow of H +, and inhibit the decrease of the concentration of H + in senescent lysosomes.
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