Molecular Aspects of Adipose-Derived Stromal Cell Senescence in a Long-Term Culture: A Potential Role of Inflammatory Pathways

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Abstract
Long-term culture of mesenchymal stromal/stem cells in vitro leads to their senescence. It is very important to define the maximal passage to which the mesenchymal stromal/stem cells maintain their regenerative properties and can be used for cellular therapies and construction of neo-organs for clinical application. Adipose-derived stromal/stem cells were isolated from porcine adipose tissue. Immunophenotype, population doubling time, viability using bromodeoxyuridine assay, MTT assay, clonogenicity, β-galactosidase activity, specific senescence-associated gene expression, apoptosis, and cell cycle of adipose-derived mesenchymal stromal/stem cells (AD-MSCs) were analyzed. All analyses were performed through 12 passages (P). Decreasing viability and proliferative potential of AD-MSCs with subsequent passages together with prolonged population doubling time were observed. Expression of β-galactosidase gradually increased after P6. Differentiation potential of AD-MSCs into adipogenic, chondrogenic, and osteogenic lineages decreased at the end of culture (P10). No changes in the cell cycle, the number of apoptotic cells and expression of specific AD-MSC markers during the long-term culture were revealed. Molecular analysis showed increased expression of genes involved in activation of inflammatory response. AD-MSCs can be cultured for in vivo applications without loss of their properties up to P6.

Keywords
adipose-derived stromal/stem cell, aging, senescence, long-term culture, gene expression

Introduction
Multipotent mesenchymal stromal/stem cells isolated from adipose tissue in last years have become the leading type of cells used for tissue engineering application. Advantage of this cell source compared to other sources like bone marrow is greater availability and minimally invasive adipose tissue harvesting technique. Isolated cells have highly proliferative and multidifferentiative potential. Properties of AD-MSCs (adipose-derived mesenchymal stromal/stem cells) were well established, thanks to increasing use of this cell type in large number of studies. Analysis of immunophenotype, proliferation and differentiation potential, clonogenity, and other tests are standard procedures performed to confirm multipotent properties of isolated cells. These assessments were performed in most studies at early stages of culture, usually after three or four passages. Data about AD-MSC characteristics during long-term culture are...
missing\textsuperscript{5,6}. Construction of large organs like neo-bladder or neo-conduit using tissue engineering techniques require large number of cells. It was indicated that cells seeded in large density (50 $\times$ 10\textsuperscript{6}/cm\textsuperscript{2}, 10 $\times$ 10\textsuperscript{6}/cm\textsuperscript{2}) promoted urinary bladder regeneration more effectively protecting from scar formation\textsuperscript{7,8}. Higher cell density increases process of regeneration by extending time of viable cells secreting growth factors, which enhance proper well-organized cell layers reconstruction\textsuperscript{8}. Such high cell number can be difficult to achieve from early passages especially in cases when the amount of adipose tissue is limited. That is why it is very important to show how long AD-MSCs retain their regenerative properties and determine the maximum passage after which AD-MSCs can be used for clinical applications without signs of senescence.

The aim of this study was to analyze the regenerative properties of AD-MSCs isolated from porcine adipose tissue in long-term culture. Immunophenotype, proliferation and differentiation potential, clonogenity, senescence markers, cell cycle, and apoptosis were analyzed in order to determine the maximum cultivation time after which AD-MSCs maintain their regenerative properties and can be used for in vivo studies.

Materials and Methods

Animals

The study was performed with the permission of the Local Ethics Committee (no. 30/2013). Ten female domestic pigs included in this study weighed between 30 and 40 kg (age 10 to 12 wk). Adipose tissue was harvested from subcutaneous tissue of the abdominal wall. Adipose tissue immediately after resection was transferred to containers with Dulbecco’s Modified Eagle’s Medium/ Ham’s F12 (GE Healthcare Life Science, Logan, UT, USA) supplemented with antibiotics: penicillin (100 U/ml, GE Healthcare Life Science), streptomycin (100 $\mu$/ml, GE Healthcare Life Science), amphotericin B (5 $\mu$/ml, BD Bioscience, Franklin Lakes, NJ, USA) and transported to the laboratory.

Isolation and Culture of Porcine Adipose-derived Stromal/Stem Cells

The adipose tissue was washed twice with deionized water and twice with phosphate-buffered saline (PBS; Pan-Biotech, Aidenbach, Germany) supplemented with antibiotics: 100 U/ml penicillin, 100 $\mu$/ml streptomycin, and 5 $\mu$/ml amphotericin B. The adipose tissue was purified by blood vessel resection. Next, 32 g of adipose tissue from each animal was washed twice with PBS supplemented with antibiotics, cut to small pieces and equally divided into four tubes. Samples were digested in the mixture of collagenase P (1 mg/ml, Sigma-Aldrich, Steinheim, Germany), calcium chloride (5 mM, Sigma-Aldrich), Hank’s Balanced Salt Solution (Pan-Biotech), and HEPES Buffer (PAA, Pasching, Austria) at a concentration of 1 ml enzyme/1 g of tissue in a shaking water bath for 60 min at 37°C. After digestion, the collagenase P was neutralized by adding an equal volume of DMEM/Ham’s F12 supplemented with 10\% fetal bovine serum (FBS; Pan-Biotech), 100 U/ml penicillin, 100 $\mu$/ml streptomycin, and 5 $\mu$/ml amphotericin B. Next, the samples were filtered through 100 $\mu$m cell strainer (BD Bioscience) and centrifuged at 1,200 $\times$ g for 5 min. After centrifugation the supernatant was removed using Pasteur pipette (BD Bioscience) and the pellet was resuspended in DMEM/F12 supplemented with 10$\%$ FBS, 10 ng/ml basic fibroblast growth factor (Thermo Fisher Scientific, Waltham, MA, USA), 100 U/ml penicillin, 100 $\mu$/ml streptomycin, and 5 $\mu$/ml amphotericin B. Number of cells was determined by trypan blue exclusion. Isolated cells were seeded at a density of 2 $\times$ 10\textsuperscript{6} cells/cm\textsuperscript{2}. Cells were cultured at 37°C in 5$\%$ CO\textsubscript{2} and 95$\%$ humidity. Growth medium was changed every 2 d. Cell morphology and growth were evaluated under inverted light microscope (Leica, Wetzlar, Germany). Cells were cultured until the 12th passage.

Cell Surface Markers Expression

Phenotypic analysis of cell surface markers, including CD29, CD44, CD90, CD11b, CD31, and CD45, was performed every second passage. Appropriate isotype controls were used to determine the levels of background fluorescence. For each marker, approximately 0.5 $\times$ 10\textsuperscript{6} cells were used for analysis. Cells were washed twice in PBS, centrifuged for 5 min at 700 $\times$ g and suspended in 200 $\mu$l of Staining Buffer (BD Bioscience) containing 2$\%$ FBS and 0.09$\%$ sodium azide. Appropriate amounts of specific antibodies were pipetted to each tube (Table 1). Cells were then incubated for 30 min in the dark at 4°C. After incubation cells were washed twice with Staining Buffer and analyzde with the use of FACSCanto II flow cytometer (BD Bioscience).

Population Doubling Time (PDT)

Growth kinetics in long-term culture was monitored by counting cell number in subsequent passages with the use of hemocytometer and trypan blue. PDT was calculated according to the formula PDT = $T \ln(N2/N1)$, where $T$ is the incubation time, $N1$ is the cell number at the beginning of the incubation, and $N2$ is the cell number at the end of the incubation.

Proliferation

Cell proliferation was examined by bromodeoxyuridine (BrdU) incorporation at each subsequent passage according to the manufacturer protocol (Merck Millipore, Burlington, MA, USA). For this purpose adipose-derived stromal/stem cells were seeded on 96-well plates (Corning, Manassas, VA, USA) at a density of 4 $\times$ 10\textsuperscript{3} cells per well. After 24-
Table 1. Antibodies Used for Analysis of the Cell Surface Marker Expression.

| Antibody                          | Concentration (µg/ml) | Distributor        |
|-----------------------------------|-----------------------|--------------------|
| Anti-Integrin beta 1 antibody     | 0.75                  | Abcam, UK          |
| [MEM-101A] (FITC)                 |                       |                    |
| Anti-CD44 antibody [MEM-263]      | 0.5                   | Abcam, UK          |
| (FITC)                            |                       |                    |
| Anti-CD90/Thy1 antibody [SE10]    | 0.5                   | Abcam, UK          |
| (FITC)                            |                       |                    |
| Anti-CD11b antibody [2F4/11]      | 0.5                   | Abcam, UK          |
| (FITC)                            |                       |                    |
| Mouse Anti-Rat CD31 Clone [TLD-3A12] (PE) | 0.5 | BD Biosciences, USA |
| CD45 antibody [K252-1E4] (FITC)   | 0.5                   | GeneTex, USA       |
| Mouse IgG1 kappa [MOPC-21] (FITC) | 0.5                   | Abcam, USA         |
| Mouse IgG1, kappa[MOPC-31C] (PE)  | 0.5                   | BD Biosciences, USA|

FITC: fluorescein isothiocyanate; PE: phycoerythrin.

h culture, 20 µl of the diluted BrdU label was added to the appropriate wells. At the end of the incubation time (48 h) cell culture medium was removed and 200 µl of Fixing Solution was added to each well. Cells were fixed for 30 min at room temperature. After fixation, cells were rinsed with 1× Wash Buffer and incubated with 100 µl of prediluted anti-BrdU monoclonal antibody for 1 h at room temperature. Then cells were again washed and incubated with goat anti-mouse immunoglobulin G peroxidase conjugate for 30 min at room temperature. After the final wash, 100 µl of TMB Peroxidase Substrate was added to each well and cells were incubated for 30 min at room temperature in the dark. To stop the reaction 100 µl of acid Stop Solution was pipetted to each well. The absorbance was measured at 450 nm (Varioskan LUX, Thermo Fisher Scientific).

Viability

Cell viability was examined by MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay at each subsequent passage. Adipose-derived stromal/stem cells were seeded on 96-well plates at a density of 4 × 10^3 cells per well. After 48 h culture, cells from each well were rinsed with PBS (Corning) and incubated with 50 µl of 1 mg/ml MTT solution (Sigma-Aldrich) for 2 h in 37°C. After the incubation, MTT solution was removed and formazan crystals were dissolved in 200 µl of dimethyl sulfoxide (POCH, Poland). The absorbance was measured at 570 nm (Varioskan LUX, Thermo Fisher Scientific).

Clonogenicity

Reproductive viability was examined by colony-forming assay every second passage. Adipose-derived stromal/stem cells were seeded on six-well plates (Corning) at a density of 1 × 10^5 cells per well. After 10-d culture, cell colonies were rinsed with 2 ml of 0.9% saline (Avantor, Gliwice, Poland) and stained with 0.25% methylene blue solution (Sigma-Aldrich). Colonies containing more than 50 individual cells were counted using a stereomicroscope (Leica).

Apoptosis

DNA fragmentation was detected with the APO-DIRECT Kit (BD Biosciences) according to the manufacturer protocol. Every second passage, approximately 1 × 10^6 cells were fixed in 1% paraformaldehyde for 30 min. Then cells were centrifuged for 5 min at 700 × g, suspended in PBS, and after subsequent centrifugation fixed in 70% ethanol (Avantor). After overnight incubation at −20°C, the cells were centrifuged and suspended in 1× Wash Buffer. In the next step cells were suspended in 50 µl of DNA Staining Solution containing terminal deoxynucleotidyl transferase and FITC-dUTP (fluorescein isothiocyanate-deoxyuridine triphosphate) for 60 min at 37°C. At the end of the incubation time 1 ml of Rinse Buffer was added and cells were centrifuged. Then cells were suspended in 1 ml of Rinse Buffer and 0.5 ml of the PI/RNase Staining Buffer. After 30 min incubation in room temperature, cells were analyzed with the use of FACSCanto II flow cytometer (BD Biosciences). Results were compared to negative (live cells) and positive (apoptotic cells) controls provided by the manufacturer.

Cell Cycle

Cell cycle progression was determined with the use of Tali Cell Cycle Kit (Thermo Fisher Scientific) followed by flow cytometry analysis. Every second passage, approximately 1 × 10^6 cells were suspended in PBS and subsequently fixed in ice-cold 70% ethanol. Cells were kept in ethanol overnight at −20°C and then centrifuged for 5 min at 700 × g. The obtained cell pellet was suspended in PBS. After subsequent centrifugation cells were suspended in 200 µl of Staining Solution composed of propidium iodide, RNase A, and Triton X-100, incubated for 30 min in the dark at room temperature and analyzed with the use of FACSCanto II flow cytometer. Obtained data were analyzed using FlowJo v10 (Becton, Dickinson and Company, USA). Percentage of cells in G0/G1, S, and G2/M phases were calculated using Dean–Jett–Fox model.

Senescence-Associated β-Galactosidase Activity

β-galactosidase activity in senescent cells was measured cytochemically at pH 6 every second passage with the use of Senescence Cells Histochemical Staining Kit (Sigma-Aldrich). Adipose-derived stromal/stem cells were seeded on 35-mm tissue culture dishes at a density of 2 × 10^5 cells per dish. After 48 h culture, cells were rinsed with PBS and fixed for 5 min at room temperature with 1.5 ml of 1× Fixing...
Buffer. Cells were then rinsed with PBS and finally 1 ml of Staining Mixture, prepared according to the manufacturer protocol, was added to each dish. Dishes were incubated overnight at 37°C without CO₂. Stained cells were counted under an inverted microscope (Leica).

**Differentiation Potential—Adipogenesis**

AD-MSCs differentiation into adipocytes was induced with the use of Mesenchymal Adipogenesis Kit (Merck Millipore). Adipose-derived stromal/stem cells at P2, P6, and P10 were seeded on six-well plates (Corning) at a density of 1,2 × 10⁵ cells per well for qualitative analysis and on 96-well plates at a density of 1.2 × 10⁴ cells per well for quantitative analysis. After 72 h culture medium was replaced with 1 ml or 200 μl Adipogenesis Induction Medium, respectively, for 12- and 96-well plates. Medium was changed every 2 d for 21 d following differentiation schedule provided by the manufacturer. Differentiated cells were fixed with 4% formaldehyde (Sigma-Aldrich) for 30 min at room temperature, washed with PBS for 10 min, and subsequently washed with distilled water. Lipid droplets were observed under an inverted microscope.

After 21-d differentiation into adipocytes, accumulation of intracellular triglycerides was determined with AdipoRed Assay Reagent (Lonza, Verviers, Belgium). After washing with PBS, 5 μl of AdipoRed was pipetted to each well. Plates were incubated for 10 min at 37°C and then fluorescence was measured with excitation at 485 nm and emission at 572 nm (Varioskan LUX, Thermo Fisher Scientific).

**Differentiation Potential—Osteogenesis**

Differentiation into osteocytes was induced using StemProOsteogenesis Differentiation Kit (LifeTechnologies, Carlsbad, CA, USA). Adipose-derived stromal/stem cells at P2, P6, and P10 were seeded on six-well plates at a density of 1.8 × 10⁴ cells per well for qualitative analysis and on 96-well plates at a density of 2 × 10³ cells per well for quantitative analysis. After 72-h culture, medium was replaced with 1 ml or 200 μl prewarmed Complete Osteogenesis Differentiation Medium, respectively, for 12- and 96-well plates. Medium was changed every 3 d for 21 d following differentiation schedule provided by the manufacturer.

After 21 d of differentiation extracellular calcium depositions were specifically stained bright orange-red using Alizarin Red S. Before staining, cells were fixed with 4% formaldehyde for 30 min at room temperature and rinsed twice with distilled water. Cells were then stained with 2% Alizarin Red S Solution (Sigma-Aldrich) for 2 min at pH 4.2. After subsequent washing with water, calcium deposits were visualized under inverted microscope.

Differentiation hydroxyapatite portion of the bone-like nodules deposited by cells was quantitated with the use of OsteoImage Assay (Lonza). Differentiated cells were rinsed with PBS and fixed with 4% formaldehyde for 30 min at room temperature. After fixation, cells were washed twice with 1× Wash Buffer and then incubated with 100 μl of Staining Reagent at room temperature, protected from the light, for 30 min. Cells were then rinsed twice with Wash Buffer. After final wash 200 μl of diluted wash buffer was added to all wells and fluorescence was measured with excitation at 492 nm and emission at 520 nm (Varioskan LUX, Thermo Fisher Scientific).

**Differentiation Potential—Chondrogenesis**

Chondrogenic differentiation of adipose-derived stromal/stem cells was induced with the use of StemProChondrogenesis Differentiation Kit (LifeTechnologies). Adipose-derived stromal/stem cells at P2, P6, and P10 were seeded on 12-well plate in 5 μl droplets containing 8 × 10⁴ cells to generate micromass cultures. After 2-h incubation 1 ml of Chondrogenesis Induction Medium was added and cultures were refeeded every 3 d following differentiation schedule provided by the manufacturer.

After 14 d of differentiation into chondrocytes, cells were fixed with 4% formaldehyde for 30 min at room temperature and washed twice with PBS. For proteoglycans visualization micromass cultures were then stained with 1% Alcian Blue (Sigma-Aldrich) for 30 min. After subsequent washing with 0.1 N HCl and in distilled water, chondrogenic pellets were visualized under inverted microscope.

**Molecular Markers of Senescence**

AD-MSCs from the P2, P6, and P10 were placed in RNA-later solution (Thermo Fisher Scientific) and stored at −80°C. RNA was isolated using RNeasy Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s protocol. Quality and quantity of RNA were evaluated using NanoDrop (Thermo Fisher Scientific) and Agilent 2100 BioAnalyzer (Agilent Technologies, Santa Clara, CA, USA). Complementary DNA was synthesized from 500 ng of total RNA using RT² First Strand Kit (Qiagen). Gene expression of senesced cells was determined by real-time quantitative reverse transcriptase polymerase chain reaction according to RT² Profiler PCR Array Handbook using RT² Profiler PCR Array Pig Aging (Qiagen) on LightCycler 480 (Roche, Basel, Switzerland) (supplemental Table S1). The data analysis was performed at GeneGlobe Data Analysis.

**Statistical Analysis**

Each experiment was performed at least in triplicate. Average cell viability was expressed as a percentage relative to
Table 2. Immunophenotypic characterization of adipose-derived mesenchymal stromal/stem cells (AD-MSCs) in subsequent passages.

| Passage | CD29 (%) | CD44 (%) | CD90 (%) | CD11b (%) | CD31 (%) | CD45 (%) |
|---------|----------|----------|----------|-----------|----------|----------|
| P2      | 36.57 ± 8.52 | 85.67 ± 10.75 | 89.69 ± 9.17 | 0.18 ± 0.20 | 10.91 ± 8.74 | 0.35 ± 0.48 |
| P4      | 57.53 ± 7.80 | 87.15 ± 9.42 | 88.18 ± 9.15 | 0.10 ± 0.00 | 8.01 ± 2.53 | 0.38 ± 0.33 |
| P6      | 21.70 ± 14.12 | 87.12 ± 15.45 | 81.03 ± 12.12 | 0.10 ± 0.00 | 12.35 ± 7.95 | 0.28 ± 0.28 |
| P8      | 24.53 ± 13.40 | 89.27 ± 11.64 | 74.98 ± 10.02 | 0.10 ± 0.00 | 12.09 ± 7.85 | 0.53 ± 0.60 |
| P10     | 49.63 ± 11.41 | 85.80 ± 11.92 | 83.18 ± 8.19 | 0.10 ± 0.00 | 6.48 ± 2.97 | 0.33 ± 0.41 |

Values presented as a mean ± standard deviation from three independent measurements.

the control. All data were presented as means ± standard deviation. Statistical analysis was performed with Student's t-test using SPSS Statistica (Predictive Solutions, Cracow, Poland). The values were compared to the data from the second passage (to the first passage in the case of cell number and PDT analysis), which served as a control. Values of P lower than 0.05 were considered as statistically significant.

**Results**

**Immunophenotype Analysis**

Cultured AD-MSCs maintained stable expression of CD44⁺ (87.00% ± 2.24), CD90⁺ (83.41 ± 1.49), CD11b⁻ (0.12% ± 0.09), and CD45⁻ (0.38% ± 0.13) in subsequent passages (Table 2). Relatively low expression of CD29⁺ ranging between 57.5% ± 7.8% at P4 and 21.7% ± 14.1% at P6 was observed. Expression of endothelial cell marker CD31 during long-term culture was slightly increased, between 12.35% ± 7.95% at P6 and 6.48% ± 2.97% at P10 (Table 2).

**Analysis of Growth Kinetics and Morphology of AD-MSCs**

To examine the long-term growth kinetics, the cell number in subsequent passages, starting with the first passage of AD-MSC primary culture, was determined. The highest average number of AD-MSCs obtained at P1 and P2, respectively, was 15.7 × 10⁶ and 16.0 × 10⁶ cells. After P2 we observed a gradual decrease in cell number (Fig. 1A). The decrease in the number of cells was accompanied by an increase in the PDT (Fig. 1B). Average PDT between P2 and P12 increased 2.8-fold. Statistically significant decrease in PDT was observed at P7 and subsequent passages (Fig. 1B).

During the long-term AD-MSC culture we observed morphological changes typical for senescent cells. At early passages AD-MSCs were morphologically a homogenous population of small and spindle-shaped cells (Fig. 2A). Between P6 and P8 the cells became much larger and got irregular shape (Fig. 2F). The appearance of dark inclusions in cytoplasm was observed from P6 (Fig. 2C).

**Analysis of AD-MSC Proliferation**

Relative levels of BrdU incorporated into DNA during the S phase of the cell cycle gradually decreased at subsequent passages (Fig. 1C). The highest proliferation index was observed at P2. Between P2 and P3, BrdU incorporation decreased significantly by 29.6% (P < 0.05). At late passages (>P10) BrdU incorporation remained relatively constant.

**Viability Analysis**

The differences in cell viability between P2 and P8, as measured spectrophotometrically, were not statistically significant (values between 0.68 ± 0.10 and 0.55 ± 0.08, P > 0.05). The highest decrease in cell viability was observed between P9 and P10, approximately 34.2%. Similar to BrdU incorporation, mitochondrial dehydrogenase activity remained constant in late passages (>P10) (Fig. 1D).

**Clonogenicity Analysis**

The colony-forming potential of AD-MSCs decreased with subsequent passages. Clonogenic efficiency between P2 and P6 ranged between 75.2 and 63.2 per 1 × 10³ cells plated (Fig. 1E). Statistically significant reduction in the AD-MSCs clonogenicity by 54.4% appeared at P8 (P < 0.05). Clonogenic potential at P10 and P12 decreased, respectively, to 23.5 and 20.6 per 1 × 10³ cells plated (Fig. 1D).

**DNA Fragmentation Analysis**

Activation of endonucleases during the apoptotic program was analyzed on the basis of DNA breaks labeling with FITC-dUTP, followed by flow cytometric analysis. No differences in fluorescence intensities at subsequent passages after staining cells with FITC-labeled anti-BrdU antibody were observed (Fig. 3C). Average fluorescence intensities of DNA breaks in AD-MSCs between P2 and P10 ranged between 0.13% and 0.55% ± 0.24% and were similar to negative control cells (0.86% ± 0.21%) (P > 0.05).

**Cell Cycle Analysis**

Analysis showed lack of changes in cell distribution between each cell cycle phase in long-term culture (Fig. 3A, B).
Fig. 1. Properties of AD-MSCs during long-term in vitro culture. (A) Average number of AD-MSCs at subsequent passages. (B) Average population doubling time of AD-MSCs at subsequent passages. (C) Cell proliferation assessed on the basis of bromodeoxyuridine incorporation into newly synthesized DNA strands. (D) Cell viability measured with the use of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide assay. (E) Clonogenic potential of AD-MSCs in subsequent passages. Bars represent standard deviation. Statistically significant differences: *P < 0.05, **P < 0.001.

AD-MSC: adipose-derived mesenchymal stromal/stem cell; OD: optical density.

Fig. 2. Morphology of adipose-derived mesenchymal stromal/stem cells during a long-term in vitro culture. Passages: P2(A), P4(B), P6(C), P8(D), P10(E), P12(F). Small and spindle-shaped cells (A); appearance of dark inclusions (C, arrow); cells become much larger and get irregular shape (F). Images were taken on the Nikon phase-contrast microscope (objective magnification 10×).
Differences were observed only in P8 in which more cells cumulate in G2/M phase, and this effect was not observed in other passages.

**Analysis of AD-MSCs’ Differentiation Potential**

Adipogenic differentiation confirmed by lipid vesicles formation was the most effective at P2 (Fig. 4A). At P6 and P10 accumulation of intracellular triglycerides in differentiated cells was similar to undifferentiated control.

Surprisingly, the most effective osteogenic differentiation was noted at P6, when 4.65-fold increase in hydroxyapatite portion of bone-like nodules deposited by cells was observed (Fig. 4A, B). Chondrogenic differentiation was confirmed on the basis of Alcian Blue staining. Despite the same initial cell number used for micromass culture generation, cell pellets from P2 were larger and denser than those from P6 and P10 (Fig. 4A). The Alcian Blue staining intensity decreased with subsequent passages.

**β-Galactosidase Activity Analysis**

At early passages (<P8) only single cells were positive for senescence-associated β-galactosidase activity (SA-β-gal). A gradual and significant increase of β-galactosidase-positive senescence cells was observed from P8 to P12 ($P < 0.05$). The number of AD-MSCs positive for SA-β-gal increased from 70.0 at P8 to 153.3 at P12 (Fig. 5).

**Molecular Analysis of Senescence-Related Genes**

Molecular analysis revealed a different senescence gene expression profile in subsequent AD-MSC passages (Fig. 6, Table 3). After P6 compared to P2 we observed overexpression of 18 genes and underexpression of 11 genes. After P10, the number of overexpressed genes increased up to 29 and underexpressed to 13 (Fig. 6A, B). The hierarchical clustering performed for all analyzed passages divided differentially expressed genes (DEGs) into two main clusters: AD-MSCs from P2 together with P6 and AD-MSCs from P10. The gene expression profile in cells at P2 was comparable to the profile observed in cells at P6. This differed from the gene expression profile observed in AD-MSCs at P10 (Fig. 6C). Analysis of DEGs using the STRING database version 11.0 allows for detection of four pathways, which could play potential role in AD-MSCs aging. The largest number of genes (eight) was involved in complement and coagulation cascade (C3, C4, C1QA, C1QC, CFH, CLU, C3AR1, C5AR1), six genes were involved in phagosome activation (C3, CTSS, CD14, TLR2, TLR4, and FCGR1A), four genes in Toll-like receptor signaling pathway (CD14, TLR2, TLR4, and SPP1), and three genes in NOD-like receptor signaling pathway (TLR4, CASP1, and TXNIP). Analysis of these genes using STRING and KEGG PATHWAY databases detected interactions between all four pathways. As a result of this analysis we concluded that senescence of AD-MSCs was related to alteration of genes’ expression involved in activation of inflammatory process leading to growth inhibition and cell death through phagocytosis (Fig. 7).

**Discussion**

In this study we analyzed the biological properties of porcine AD-MSCs in a long term in vitro culture (12 passages) focusing on molecular aspects of cellular senescence. We
Fig. 4. Differentiation potential of adipose-derived mesenchymal stromal/stem cells during long-term culture. (A) Experiment assessed on the basis of qualitative analysis of lipid vesicles formation (adipogenesis using Oil Red O staining), presence of mineralized nodules (osteogenesis using Alizarin Red staining) and proteoglycans synthesis (chondrogenesis using Alcian Blue staining). (B) Experiment using quantitative analysis using AdipoRed Assay Reagent (Lonza) for adipogenesis and OsteoImage Assay (Lonza) for osteogenesis. Results obtained in P6 and P10 were statistically significant compared to P2 ($P < 0.01$).

|          | Adipogenesis | Osteogenesis |
|----------|--------------|--------------|
| P2       | 1.60±0.11    | 3.74±0.36    |
| P6       | 1.04±0.02    | 4.65±0.52    |
| P10      | 0.99±0.08    | 0.90±0.14    |

Fig. 5. β-galactosidase expression in long-term AD-MSC culture. (A) Number of SA-β-gal-positive AD-MSCs at different passages. (B) AD-MSCs stained for SA-β-gal. Arrows indicate senescent cells. Scale bar, 200 μm. Statistically significant differences: **$P < 0.001$.

AD-MSC: adipose-derived mesenchymal stromal/stem cell.
cultured AD-MSCs till P12 because after that time (about 40 d of culture) they changed significantly their morphology and decreased proliferation rate what in consequence leads to getting even smaller cell numbers than initially seeded. Obtained results showed that during the first eight passages most of the AD-MSC properties were on the same comparable to P2 level. Only significant decrease in cell number and BrdU incorporation after P3 and P2, respectively, was observed (Fig. 1A, C). After P8 we observed about 50% reduction in cell number compared to P2 what can be a potential problem in obtaining a suitable cell number for cell therapies or construction of neo-tissues or neo-organs using tissue engineering approach. Doubling times ranged between 30 and 50 h till P10 and increased up to 90 h at P12, which is consistent with another work in which dental pulp and bone marrow mesenchymal stem cells were analyzed. In another study conducted on human and rhesus bone marrow mesenchymal stem cells (BM-MSCs) and AD-MSCs, doubling time reach 120 h at the end of culture (P20–P30). Mesenchymal stromal cells isolated from human chorionic villi kept the doubling time value on the level of 30 h till P10, reaching about 60 h at P20. Another study performed on BM-MSCs together with umbilical cord mesenchymal stem cells (UC-MSCs) also showed decreasing proliferation capacity with prolonged culture time (P12).

Morphological changes typical for senescent cells were observed after the P6 (about 21 d). Similar findings were noticed for BM-MSCs. After 84 d of culture intracellular granules were observed. Longer culture resulted in cell vacuolization and cell fragmentation visualized as debris in culture medium. In another study performed on human AD-MSCs the first morphological changes appeared after the P7, with the cells becoming larger with wider cytoplasmatic projections. Higher passages (P20–P30) resulted in irregular morphology with more granular cytoplasm and increasing number of cellular debris in culture medium. During subsequent passages shortening of telomeres was observed. In our study cell senescence was analyzed by measuring β-galactosidase activity and analysis of senescence-related gene expression. Increasing expression of β-galactosidase appears in aging cells, which is a consequence of morphological changes with the increase in the abundance of lysosomal enzyme, which could be linked with increased lysosomal biogenesis observed in senescent cells. In our study, level of β-galactosidase activity increased gradually with subsequent passages (Fig. 5), which was compatible with changes in cell morphology (Fig. 2). Similar results were observed in hair follicle mesenchymal stem cells (HF-MSCs) in which aging cells in greater number started to appear after P8 (in our study after P6). In the case of human and rhesus BM-MSCs and AD-MSCs

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**Fig. 6.** Changes in senescence-related gene expression in adipose-derived mesenchymal stromal/stem cell culture. (A, B) Results from RT² Profiler PCR Arrays Pig Aging plate. (C) Hierarchical clustering of differentially expressed genes.
significant increase in the number of aging cells (76% and 95%, respectively) appeared after P20, compared to P12 in our study. Another symptom indicating cell senescence is loss of differentiation potential. We observed decrease of adipogenic and chondrogenic potential after P6 and osteogenic potential after P10. In the study of Bonab et al., decrease of BM-MSCs’ osteogenic potential after P8 and adipogenic after P6 was observed. Similar observation was done in the case of HF-MSCs, and difference was noticed only in the case of osteogenic potential, which increased after P8 compared to P5 but significantly decreased after P11. Human and rhesus BM-MSCs and AD-MSCs retained their multipotential capacity at least up to P10. Study conducted on omentum AD-MSCs showed that even long-term culture (P20) did not affect osteogenic and adipogenic potential. In another study even after P15 no differences in adipogenic, chondrogenic, and osteogenic potential of tonsil MSCs was observed, and long-term culture effected in decrease of proliferative potential measured by colony-forming unit and MTT assay. Colony-forming unit decreased about 35% at P15 passage (in our study about 54% after P8).

Changes in cell cycle profile were observed in dental pump, peripheral blood, and umbilical cord blood MSCs. Additionally, in the case of peripheral blood mesenchymal stem cells (PB-MSCs) and UC-MSCs increase in apoptotic cell number was noticed. Increase in apoptotic cell number appeared after P20 in the case of UC-MSCs. In our study no changes in cell distribution in cell cycle phases and in number of apoptotic cells were observed throughout the study period (Fig. 3C).

Mesenchymal stromal/stem cells are characterized by expression of specific surface markers like CD90, CD44, or CD29 and lack of antigen expression like CD11b, CD31, and CD45. Our results showed that expression of most studied markers was stable in long-term culture, which is consistent with the results of Marappagounder et al. who also noticed maintenance of characteristic AD-MSC markers in prolonged culture. Another study conducted on human AD-MSCs also confirmed stability of their phenotype in long culture, even after P30. Similar results were also observed in the case of BM-MSCs and UC-MSCs. In our study instability of two antigens, CD29 and CD31, was observed. According to the recommendation of the International Society for Cell & Gene Therapy both CD29 and CD31 are not considered in minimal criteria to define multipotent MSCs; however, in literature these two antigens were examined for identification of AD-MSCs. Expression of surface markers can vary between MSCs obtained from different sources, for example in the case of AD-MSCs positive expression of CD34 can be detected while in general this is a negative marker in other types of MSCs. According to this information positive expression of CD31 antigen in the case of AD-MSCs is also possible. Criteria for MSC identification were established for human cells. We have performed experiment on porcine cells, which probably could affect final results.

From our knowledge this is the first study in which porcine AD-MSCs were analyzed for senescence genes expression using RT2 Profiler PCR Array. Molecular analysis showed a different expression pattern of genes involved in senescence. After P6 compared to P2 we observed overexpression of 18 genes and underexpression of 11 genes. After P10 the number of overexpressed genes increased up to 29 and underexpressed to 13. Analysis of differentially expressed genes using the STRING database version 11.0 allows for identification of four pathways which could play a potential role in AD-MSCs aging: complement and

| Expression | P6 vs. P2 | P10 vs. P2 |
|------------|-----------|-----------|
| Upregulated | ANGEL2 | ANGEL2 |
| | ANXA5 | ANXA3 |
| | ARL6IP6 | C15 |
| | C3AR1 | C3 |
| | CA4 | C3AR1 |
| | CALB1 | C4A |
| | CD14 | CSAR1 |
| | COL3A1 | CA4 |
| | CTSS | CALB1 |
| | CX3CL1 | CASP |
| | LMNA | CLU |
| | LYZ | COL3A1 |
| | MBP | CTSS |
| | POT1 | CX3CL1 |
| | SIRT6 | EP300 |
| | TERF1 | FGR1 |
| | TMEM33 | FOXO1 |
| | TXNIP | JKMIP3 |
| | LOC100628185 | |
| | LYZ | MBP |
| | RNF144B | TLR2 |
| | TLR4 | TPP1 |
| | TXNIP | TXNIP |
| | VPS13C | VLC |
| | VWA5A | ZNF25 |

| Downregulated | APOD | APOD |
| | C1QA | C1QC |
| | C1QC | CFH |
| | C3 | FER1G |
| | C4A | FCR3B |
| | CFH | LMNB1 |
| | FCR1G | LMNB2 |
| | FCR3B | LOC102164975 |
| | S100AB | LSM5 |
| | S100A9 | S100A9 |
| | SPP1 | SPP1 |
| | TFAM | |

Table 3. Differential Senescence-Related Gene Expression in Subsequent Passages.
coagulation cascade, phagosome, Toll-like receptor signaling pathway, and NOD-like receptor signaling pathway. These pathways are activated mainly as a result of immunological response. Senescent cells are regarded as proinflammatory and immunogenic, playing active roles both in activation and recruitment of the immune system. Additionally senescent cells may persist indefinitely within tissue until their removal by the immune system, necrosis, or other forms of cell death. All four pathways detected in this study are directly connected to each other. A pathway that connects all pathways together seems to be the Toll-like receptor signaling pathway. In our study we observed overexpression of four genes involved in this pathway (TLR2, TLR4, CD14, and SPP1) from which TLR4 seems to play a key role. This protein is also involved in NOD-like signaling receptor pathway leading to activation of caspase 1, which is additionally activated by NLRP3 (activated also by TXNIP overexpressed in our study). Activation of Caspase-1 induces maturation and releases proinflammatory cytokines (IL18 and IL1β). Creation of NLRP3 inflammasome is closely related to cellular senescence and was observed in endothelial cells, cardiac fibroblasts, and primary BM-MSCs. Toll-like signaling pathway is involved in activation of complement and coagulation cascade; in our study, five genes, involved in the complement cascade pathway, were overexpressed after P12 (C3, C4A, CLU, C3AR1, and C5AR1). C3 protein leads to activation of CR1 and CR3 genes, which consequently induce phagocytosis, and this process is also activated by overexpression of Toll-like receptor signaling pathway (CD14, TLR2, and TLR4). Activation of C3AR1 and C5AR1 generates chemotaxis of phagocytic cells to the place of ongoing inflammatory process, which helps in one complement cascade function: elimination of modified or damaged host cells. Taking together, the probable mechanism of AD-MSC senescence on the basis of molecular analysis of cells after P10 is activation of immune response and phagocytosis of senescent cells (Fig. 7). Activation of immune system by senescence cells was also described by Burton and Stolzing, and cells like macrophages have the potential to cell death induction in these cells by phagocytosis. Similar mechanism of MSCs senescence was described by Shi et al., and in their study TLR4 activation was induced by S100A9. In our study both S100A9 and S100A8 were underexpressed, which indicate that expression of these proteins is not required for activation of proposed AD-MSCs’ senescence mechanism.

MSCs changed their properties (differentiation potential, growth dynamics, and cell morphology) in in vitro culture during subsequent passages. According to the obtained results, this effect is not related with donor because tissues for cell isolations were collected from animals at the same age. Observed cellular senescence can be related to medium effect. Growth conditions and metabolic needs differ significantly in vitro culture. Culture medium contains less number of growth factors and other supplements like amino acids than are normally found in extracellular tissue fluid. Additionally after isolation and during the culture cells continuously undergo proliferation, which generate unnatural stress to

Fig. 7. Potential pathway leading to phagocytosis of senescent AD-MSCs. Molecular analysis showed that during a long-term culture of AD-MSCs activation of genes involved in immune response was observed, which in consequence leads to phagocytosis of senescent cells. Overexpressed genes are marked in red. AD-MSC: adipose-derived mesenchymal stromal/stem cell.
cells. Our results showed that the best time point up to which AD-MSCs should be used for further in vitro and in vivo experiments is the P6, until which most of the AD-MSC properties were maintained comparable to early P2 level. However, after that time 50% reduction of cell number compared to first culture days was observed, which can be a potential problem in obtaining a suitable cell number for reconstruction of neo-tissues or neo-organs. The balance between proper cell number and maximum passage number necessary to obtain suitable cell amount should be preserved.

Ethical Approval
Ethical approval for this case was obtained from the Local Ethics Committee in Bydgoszcz, Poland (no. 30/2013).

Statement of Human and Animal Rights
All procedures in this study were conducted in accordance with the Local Ethics Committee in Bydgoszcz, Poland (no. 30/2013) approved protocols.

Statement of Informed Consent
There are no human subjects in this article and informed consent is not applicable.

Declaration of Conflicting Interests
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