Human Adipose-Derived Mesenchymal Stromal Cells Exhibit High HLA-DR Levels and Altered Cellular Characteristics under a Xeno-free and Serum-free Condition

Phuong T. M. Dam 1 · Van T. Hoang 2 · Hue Thi Hong Bui 1 · Le Minh Hang 1 · Duc M. Hoang 2 · Hoang Phuong Nguyen 2 · Ha Thi Lien 1 · Huong Thi Thanh Tran 1 · Xuan-Hung Nguyen 1 · Liem Nguyen Thanh 2

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Abstract

Background We have observed an increased expression of negative markers in some clinical-grade, xeno- and serum-free cultured adipose-derived mesenchymal stem/stromal cell (ADMSC) samples. It gave rise to concern that xeno- and serum-free conditions might have unexpected effects on human ADMSCs. This study aims to test this hypothesis for two xeno- and serum-free media, PowerStem MSC1 media (PS) and StemMACS MSC Expansion Media (SM), that support the in vitro expansion of ADMSCs.

Methods We investigated the expression of negative markers in 42 clinical-grade ADMSC samples expanded in PS. Next, we cultured ADMSCs from seven donors in PS and SM and examined their growth and colony-forming ability, surface marker expression, differentiation, cell cycle and senescence, as well as genetic stability of two passages representing an early and late passage for therapeutic MSCs.

Results 15 of 42 clinical-grade PS-expanded ADMSC samples showed an increased expression of negative markers ranging from 2.73% to 34.24%, which positively correlated with the age of donors. This rise of negative markers was related to an upregulation of Human Leukocyte Antigen – DR (HLA-DR). In addition, the PS-cultured cells presented decreased growth ability, lower frequencies of cells in S/G2/M phases, and increased ß-galactosidase activity in passage 7 suggesting their senescent feature compared to those grown in SM. Although MSCs of both PS and SM cultures were capable of multilineage differentiation, the PS-cultured cells demonstrated chromosomal abnormalities in passage 7 compared to the normal karyotype of their SM counterparts.

Conclusions These findings suggest that the SM media is more suitable for the expansion of therapeutic ADMSCs than PS. The study also hints a change of ADMSC features at more advanced passages and with increased donor’s age. Thus, it emphasizes the necessity to cover these aspects in the quality control of therapeutic MSC products.

Keywords Adipose-derived stem cells · Adipose-derived mesenchymal stem cells · Adipose-derived mesenchymal stromal cells · HLA-DR · Karyotype · Senescence

Introduction

Adipose-derived mesenchymal stem cells/stromal cells (ADMSCs) have great potential for cell-based therapy and regenerative medicine due to their capability of immunomodulation and in vitro differentiation into various cell lines such as adipocytes, chondrocytes, osteoblasts [1], neuronal cells [2, 3] and hepatocytes [4]. Additionally, growth factors, cytokines, exosomes and microvesicles secreted by MSCs also contribute to their regenerative activity in vivo [5–7]. Adipose tissue is an abundant source for MSCs in many clinical trials [8]. Compared to bone marrow-derived
MSCs (BM-MSCs), the harvesting procedure of ADMSCs is less invasive and can be performed through multiple sites resulting in high stem cell yield [9]. ADMSCs have been applied to treat Crohn’s fistula in a phase III trial [10], osteoarthritis and degenerative eye diseases in phase II studies [11–13]. Of note, all these trials required an in vitro expansion of isolated ADMSCs to obtain sufficient cell doses.

Although there are several commercially serum-free and xeno-free media, the expansion of ADMSCs and their biological properties in these media remain elusive. MSCs might lose their growth and multipotency upon the in vitro culture [14, 15]. Indeed, ADMSC proliferation declined after five passages in a commercial xeno-free and serum-free media called StemPro (Invitrogen, Carlsbad, CA, USA) [16]. Culture of dental pulp stem cells and alveolar bone marrow MSCs in StemPro and StemMacs_MSC/XF (Miltenyi Biotec GmbHI, Germany) resulted in reduced chondrogenic and adipogenic differentiation potential [17]. Furthermore, cardioprotective effects of MSCs diminished in a rat model for ischemia–reperfusion injury when cells of advanced passages (passage 5 to passage 10) were administrated [18]. Another challenge is that genetic changes such as tetrasyom and other chromosomal abnormalities might occur in MSC cultures [19, 20]. These can be related to spontaneous tumorigenic transformation and become a potential risk for MSC-based therapy [21]. In addition to in vitro related dysfunctions of MSCs, donor features also play an essential role in the biological activities of these cells. MSCs obtained from aged donors were associated with lower growth kinetics, increased senescence, and altered differentiation potential [22–24]. Metabolic diseases such as type 2 diabetes may restrict MSC functions [25, 26]. Therefore, the influence of culture conditions and donor characteristics on MSC characteristics should be addressed.

According to the guidance of the International Society for Cellular Therapy (ISCT), one of the criteria to define MSCs are the surface expression of CD73, CD90, CD105, CD44 and the absence of CD45, CD34, CD14/CD11b, CD19/CD79α, CD31, as well as the human leukocyte antigen HLA-DR [27, 28]. The last marker, HLA-DR, belongs to the major histocompatibility complex class II (MHC class II) and plays a crucial role in host immune responses. Allogenic MSCs were rejected in MHC class I- and MHC class II-mismatch recipient mice [29] and induce an immune response in pigs and horses [30–32]. Moreover, HLAs were demonstrated to induce the immunogenicity of allogenic ADMSCs [33]. This suggested that HLA-DR expression might induce an immune response and increase the risk of rejection in an allogeneic environment. For this reason, HLA-DR expression in cultured ADMSCs requires a special attention.

This study analyzed the immunophenotype of clinical-grade ADMSCs in a histological cohort, which were expanded under a serum-free and xeno-free condition using PowerStem MSC1 media (PS). Next, we compared the cellular properties of ADMSCs cultured in PS with their counterparts grown in our new candidate for serum-free and xeno-free medium called StemMACS MSC Expansion Media (SM).

### Materials and Methods

#### Patient Samples

The adipose tissues were obtained at Vinmec International Hospital between 2018 and 2019 and cryopreserved in the biobank after the donors signed the informed consent form. The expression of surface markers was retrospectively analyzed on 42 ADMSC samples isolated in PS (Pan-Biotech, Germany), of which 31 patients participated in a previous study [34]. This study enrolled patients with sexual functional deficiency. The other 11 samples were obtained from healthy donors. We performed a prospective study on seven samples, of which donors agreed to donate their biological materials for the current research, to investigate the cellular properties of ADMSCs cultured in different xeno-free and serum-free MSC expansion media (Suppl. Figure 1). Sample collection and data analysis were approved by the Ethics Committee of Vinmec Healthcare System and carried out in accordance with the Declaration of Helsinki.

#### Isolation, Culture, and Cryopreservation of ADMSCs

ADMSCs were isolated as previously described [35]. Briefly, adipose tissues were harvested from the donor’s lower abdomen under general anesthesia. Then adipose tissue was minced into small pieces and digested using 200 unit/ml Collagenase type I solution (Gibco, Denmark) supplemented with 200 g/l Human Albumin (Baxter, USA) at 37 °C with agitation 300 rpm/min for 60 min. The mixture was centrifuged at 1500 rpm/min for 10 min to collect the stromal vascular fraction in the pellet. Finally, the cells were resuspended in PS (Pan-Biotech, Germany) supplemented with 1% Penicillin/Streptomycin (Gibco, USA). Cells were incubated at 37 °C and 5% CO2 in a humidified atmosphere with medium exchange every three days. When reached 80–90% confluence, the cells were harvested using TrypLE™ Select Enzyme (Gibco, USA) and counted after staining with trypan blue (Life Technologies, USA).

Cells at passage 0 were cryopreserved in the Cryostor solution (Stemcell Technologies, Canada). All the cryovials were kept at -80 °C for no more than seven days and transferred to BioStore™ III Cryo (Brooks Life Sciences, USA) for long-term storage. After four months of cryopreservation, the cells were thawed rapidly at 37 °C and maintained...
in PS or SM (Miltenyi Biotec, Germany) to evaluate cell growth, phenotype, and functional properties.

**Growth Curve**

To determine the proliferative capability of ADMSCs, cells were seeded with the density of 5,000 cells/cm² in T25 flasks (Nunc) and cultured up to passage 7 as previously described [35]. Experiments were performed in triplicate for each donor. Population doubling (PD) was calculated using the formula: 

\[ PD = \frac{3.32 \times \log (a/b)}{a} \]

where \( a \) is the number of harvested cells and \( b \) is the seeding density. Finally, the population doubling time (PDT) was calculated as \( t/n \), where \( t \) is the days (or hours) of culture, \( n \) is the number of PD reached during a passage. The cell morphology was visualized under an invert phase-contrast microscope (Eclipse Ti-S/DS-Fi2-L3, Japan).

**Colony-Forming Unit (CFU) Assay**

Cells were seeded in triplicate in a six-well dish at the concentration of 20 cells/cm² and cultured in the respective media (PS or SM) for 10 to 14 days as previously described [35]. Cells were fixed in Methanol (Merck, Germany) and stained with Giemsa solution (Merck, Germany), both for 5 min at room temperature. Colonies formed on each dish were counted.

**Flow Cytometry Analysis**

ADMSCs harvested from passage 3 and 7 were assessed for surface marker expression by flow cytometry. Cells were resuspended in PBS (Gibco, USA) supplemented with 1% human serum (Pan-Biotech, Germany). The cells were incubated using the Human MSC Analysis kit (Becton Dickinson, USA) according to the manufacturer’s instruction. To examine the expression of negative surface markers individually, cells were stained with the following antibodies: CD34 Monoclonal Antibody (581) FITC (Invitrogen, USA), CD11b Monoclonal Antibody (ICRF44) PE (Invitrogen, USA), CD19 Monoclonal Antibody (HB19) PerCP-Cyanine 5.5 (Invitrogen, USA), CD45 Monoclonal Antibody (HI30) APC (Invitrogen, USA), Anti HLA-DR-FITC (Beckman Coulter, France). The samples were analyzed using BD Cantor II (Becton Dickinson, USA). At least 20,000 cells per run were acquired. The data analysis was performed using the FlowJo software (Becton Dickinson, USA).

To check viability, cells were suspended in PBS (Gibco, USA) supplemented with 1% human serum (Pan-Biotech, Germany) and stained with 7-AAD Staining Solution (Miltenyi, Germany) following the manufacturer’s instruction. For the cell cycle analysis, cells were fixed with eBioscience™ Intracellular Fixation & Permeabilization Buffer and stained with 7-AAD Staining Solution (Miltenyi, Germany) as previously described [36]. At least 20,000 cells were analyzed per run using BD Cantor II (Becton Dickinson, USA) and the FlowJo software (Becton Dickinson, USA).

**MSCs Differentiation Assays**

MSCs were differentiated into adipogenic, chondrogenic and osteogenic lineages using the StemPro® Adipogenesis Differentiation Kit, StemPro™ Chondrogenesis Differentiation Kit, StemPro™ Osteogenesis Differentiation Kit (Gibco, USA) in duplicates as previously described [35]. For this purpose, cells were cultured in either differentiation media or MSC culture media as the control for 14 days. They were then fixed by using 4% PFA (Merck, Germany) for 30 min before staining with the Oil Red O (Merck, Germany) solution to detect lipid droplets, the Alcian Blue (Sigma Aldrich) solution to find the presence of proteoglycan and Alizarin Red S (Merck, Germany) solution to mark calcium deposition. The staining was observed under an inverted microscope equipped with a camera (Olympus Corporation, Tokyo, Japan).

To perform quantitative analysis, ADMSCs were cultured in the differentiation media for 14 days. Cells of the same passage were used as the undifferentiated control. Total RNA was extracted using RNeasy Mini Kit (Qiagen, Germany) according to the manufacturer’s instruction. cDNA synthesis was carried out using the SuperScript™ IV First-Strand Synthesis System (Invitrogen/Thermo Fisher Scientific, USA), and qPCR was performed using the 7500 Fast Real-Time PCR System and PerfeCTa SYBR Green SuperMix according to the manufacturer’s instructions. The expression of the following genes was analyzed: PPARγ and Leptin for adipogenesis, ALP and PTH-R for osteogenesis, SOX9 for chondrogenesis and GAPDH for the reference gene. Primer sequences are presented in the supplementary information.

**Senescence Assay**

ADMSCs were seeded on six-well plates and cultured in the PS or SM media to obtain passage 3 and passage 7 cultures. Cell senescence was analyzed using Senescence Cells Histochemical Staining Kit (Sigma Aldrich, USA) following the manufacturer’s instruction as previously described [37]. Briefly, cells were fixed and stained with Staining Mixture containing X-gal substrate overnight after reaching 70–80% confluence. Senescent cells with β-galactosidase activity were stained blue. The total number of cells was examined using DAPI (Abcam, UK) counting. Cell images were observed and captured by a IX73 microscope and cellSens software (Olympus, Japan). Cell counting was performed using Image J software (version 1.46r). The percentage of...
senescent cells was calculated by dividing the total number of DAPI-stained cells by the number of blue-stained cells.

Karyotype

To evaluate the karyotype of ADMSCs after prolonged culture in vitro, passage 3 and 7 cells were checked for karyotype as previously described [26]. When the cells reached 80% confluence, they were incubated in KaryoMAX® Colcemid™ 10 µl/ml (Gibco, USA), diluted to 1:1000, at 37 °C in a 5% CO2 incubator for 30 min. Cells were detached from the flask by incubating in 2 ml of Trypsin EDTA 0.25% (Gibco, USA) at 37 °C within 1 to 2 min and then in a prewarmed hypotonic solution (KCl 0.56%) for 30 min. Cells were washed three times with Carnoy’s solution (3:1 v/v absolute ethanol: glacial acetic acid) before being dropped on 3–4 clean slides to spread metaphase. The slides were incubated at 60 °C overnight and stained the day after using G-banding technique. Metaphase was captured by MetaSystems, Carl Zeizz and analyzed by Ikaros software. Twelve to twenty metaphase spreads were analyzed for each sample by an experienced technician and validated by a medical genetic scientist. Karyogram is 400 band according to ISCN (An International System for Human Cytogenomic Nomenclature) 2016 standard [38].

Statistical Analysis

All experiments in this study were performed in triplicates if not otherwise indicated. Data are present as mean ± SD if not otherwise indicated. Student t-test was used for a normally distributed population, while Mann–Whitney U test was employed in nonparametric distribution. A p-value below 0.05 was considered statistically significant. Differences among multiple groups were statistically analyzed using one-way ANOVA and Tukey’s multiple comparisons test. Univariate analyses were conducted using Spearman’s rank correlation and Kruskal–Wallis test. Statistical analyses were performed using the R software version 3.6.1 by a statistician [39]. Plots were prepared using the software GraphPad Prism 5.0 (GraphPad Software, La Jolla, CA).

Results

Increased Expression of Negative Markers in ADMSC Cultures

The marker expression of ADMSCs isolated from 42 donors using the PS medium was analyzed for the MSC positive markers CD73, CD90 and CD105 and the negative or moderately expressed markers CD11b, CD19, CD34, CD45, and HLA-DR according to the ISCT’s recommendations. The samples highly expressed the markers CD73, CD90, and CD105 with more than 95% positivity (Fig. 1A and B). There are samples showing increased levels of the negative marker cocktail including CD34, CD45, CD11b, CD19, and HLA-DR (Fig. 1A). The mean values of negative marker expressing cells were 4.10% (from 0.10% to 34.24%) (Fig. 1B). Among them, 15 of the analyzed 42 samples showed more than 2% positivity for negative markers, which was the cut-off value as suggested by the ISCT. These negative marker-positive cells co-expressed CD73, CD90, and CD105 (Fig. 1A). It indicates that they represented a population of ADMSCs. Thus, our data suggest that PS medium might induce the expression of the negative markers in ADMSCs.

Next, we analyzed whether the increased expression of negative markers depends on patient characteristics such as age, gender, body mass index (BMI), and due our choice of patient cohort, of which there were 11 healthy donor and 31 patients with sexual functional deficiency. Suppl. Table 1 depicts the patient characteristics. We observed that elderly donors tended to have more negative marker-expressing cells (p = 0.035) (Fig. 1C). The other factors did not show any correlation (Suppl. Figure 2).

Elevated Expression of the Negative Markers in PS-Cultured ADMSCs caused by HLA-DR

To investigate which negative markers are increasingly expressed in the PS-cultured ADMSCs, we quantified their expression individually by flow cytometry (n = 7). For comparison, we analyzed cells derived from the same donors that were cultured in the xeno-free and serum-free SM (Fig. 2A and suppl. Figure 3). ADMSC-expanded in PS upregulated HLA-DR expression in passage 3 (mean ± SD: 27.1 ± 26.2%) and passage 7 (mean ± SD: 6.63 ± 6.59%) (Fig. 2B). In contrast, HLA-DR marker was expressed lower in ADMSC-expanded in SM, whose frequency of HLA-DR expressing cells were 0.07 ± 0.09% in passage 3 (PS versus SM: p = 0.003) and 0.03 ± 0.04% in passage 7 (p = 0.66), respectively. Although ADMSCs in the PS medium remarkably increased their HLA-DR expression in passage 3, it was decreased in the higher passage (p = 0.04). The other negative markers were absent in both cultures. Furthermore, ADMSCs grown in PS showed lower expression levels of CD90 and CD105 in passage 7 compared to passage 3 (p = 0.04 and p = 0.009, respectively) as well as those cultured in SM (p = 0.03 and p = 0.01, respectively) (Fig. 2B).

The Effect of Culture Media on Cell Growth of ADMSCs

To determine the effect of media on the cell growth, ADMSCs were cultured up to passage 7 (n = 7). ADMSCs
expanded in SM were small, spindle-shaped and could reach confluent during the analyzed period. In contrast, ADM-SCs expanded in PS also showed small size and spindle-shaped cells in early passages but became bigger and failed to reach confluent in higher passages (Fig. 3A). Accordingly, cell proliferation rate in PS was lower than that of SM in passages 6 and 7 ($p=0.04$ in both passages) (Fig. 3B). Furthermore, the frequencies of CFUs of the cells in PS were $4.31 \pm 1.13\%$ in passage 3 and $0.69 \pm 0.61\%$ in passage 7. They were significantly lower than the ones in SM, which were $11.03 \pm 2.85\%$ in passage 3 ($p<0.001$) and $9.00 \pm 3.40\%$ in passage 7 ($p<0.001$). While the CFUs of PS-expanded ADMSCs decreased in the later passage ($p=0.03$), the colony numbers of SM-cultured cells remained high in both passages ($p=0.37$) (Fig. 3C).

**Differentiation Capacity of ADMSCs**

To test the multipotency of the ADMSC cultures, the cells were differentiated into adipogenic, osteogenic,
and chondrogenic lineages. ADMSCs in both media were able to differentiate into the three cell lineages as indicated by positive staining of Oil Red O (Fig. 3D), Alizarin Red S (Fig. 3E), and Alcian Blue (Fig. 3F). Accordingly, expression of differentiation markers including PPARγ and Leptin for adipogenesis (Fig. 3G), PTH-R and ALP for osteogenesis (Fig. 3H), and SOX9 for chondrogenesis (Fig. 3I) were increased in the differentiated samples compared to their control counterparts. The expression level of PPARγ was lower in the PS cultures ($p = 0.002$), while PTH-R was expressed higher in this condition compared to the cells in the SM media ($p = 0.009$). There were no differences in the expression of the other analyzed genes.

Fig. 2 Increased HLA-DR expression in PS-cultured ADMSCs compared to their counterparts in SM. (A) A representative ADMSC sample cultured in PS and SM media and analyzed for their expression of individual negative markers using flow cytometry. (B) Differential expression of surface markers in the PS and SM cultures ($n = 7$). The cells in PS expressed increasingly HLA-DR compared to those in SM. In addition, the PS-cultured cells in passage 7 showed a decreased expression of CD90 and CD105. Levels of significance: *, $p < 0.05$. **, $p < 0.01$. Abbreviations: adipose derived mesenchymal stem/stromal cells (ADMSCs), PowerStem MSC1 media (PS), StemMACS MSC Expansion Media (SM)

Fig. 3 Morphology, cell growth, and differentiation of ADMSCs cultured in the PS and SM media. (A) Representative cell morphologies of ADMSCs cultured in the tested media from passage 1 to passage 7 ($n = 7$). (B) Population doubling time was calculated for each passage of these cells. (C) Colony-forming potential of ADMSCs in the media of interest. (D-F) ADMSCs were differentiated into adipogenic (D), osteogenic (E), and chondrogenic lineages (F) and stained with Oil Red O, Alizarin Red S, and Alcian Blue, respectively. (G-I) Quantitative gene expression analysis was performed for adipogenic markers: PPARγ and Leptin (G), osteogenic markers: PTH-R and ALP (H), and chondrogenic marker: SOX9 ($n = 6$) (I). Gray stars indicate the statistical significance comparing the differentiated cells with their undifferentiated controls. Black stars represent the statistic test between PS and SM cultures. Scale bars represent 100 µm. Levels of significance: *, $p < 0.05$. **, $p < 0.01$. ****, $p < 0.0001$. Abbreviations: adipose derived mesenchymal stem/stromal cells (ADMSCs), PowerStem MSC1 media (PS), StemMACS MSC Expansion Media (SM)
Cell Viability, Proliferation, and Senescence of ADMSCs

Cell viability of ADMSCs in PS and SM cultures were analyzed using trypan blue and 7-AAD. There were no differences in the numbers of dead cells in both passages 3 and 7 (Suppl. Figure 4). Furthermore, the cells were fixed and stained with 7-AAD to analyze their cell cycle (Fig. 4A). Apoptotic cells, which have reduced DNA content compared to living cells, were observed in the sub G1 phase. Their frequencies were increased in the PS compared to the SM cultures in both passages 3 and 7 (Fig. 4B). Accordingly, the proportion of proliferating cells in the S/G2/M phases decreased in the PS cultures in passage 7 compared to their counterparts in SM as well as the PS cells in passage 3 (Fig. 4C). The results indicated a reduced proliferation capacity of the PS cells in a higher passage.

Therefore, senescence analysis was performed using the b-galactosidase assay (Fig. 4D). Both cultures in passage 3 were free of b-galactosidase positive cells. In passage 7, cells cultured in PS showed a higher frequency of senescent cells compared to those in SM (Fig. 4E). Thus, our results of cell cycle and cellular senescence indicated that the later media was able to maintain ADMSCs longer in a proliferative state.

Changes of Karyotype of ADMSCs during in vitro Culture

To investigate whether the changes in cell growth rates, morphological and immunophenotypic characteristics as well as senescence were associated with genome instability, we performed karyotyping of ADMSCs (Fig. 5A and B). MSCs of sample 1 displayed a normal karyotype in passage 3 in both media; however, four and eight of 13 analyzed cells
### A

| Sample no. | Gender | Age | PS Passage 3 | PS Passage 7 | SM Passage 3 | SM Passage 7 |
|------------|--------|-----|--------------|--------------|--------------|--------------|
| 1          | M      | 66  | 46,XY        | 44, X,-13[4]/45, X[8]/46, XY[2] | 46,XY        | 45, XY, chtb(X)(q21),-8[1]/46, XY[19] |
| 2          | M      | 57  | Too few metaphases | Too few metaphases | 46, XY       | 46, XY       |
| 3          | M      | 48  | 46, XY       | Too few metaphases | 46, XY       | 46, XY       |
| 4          | F      | 46  | 46, XX       | Too few metaphases | 46, XX       | 46, XX       |
| 5          | F      | 43  | 46, XX       | Too few metaphases | 46, XX       | 46, XX       |
| 6          | F      | 42  | 46, XX       | 46, XX        | 46, XX       | 46, XX       |
| 7          | F      | 40  | 46, XX       | 47, XX,+2[10]/46, XX[10] | 46, XX       | 46, XX       |

### B

- **44,X,-13**
- **47,XX,+2**
- **45,XY,-8, chtb(X)(q21)**
- **46,XY**
expanded in PS at passage 7 displayed deletions of chromosome 13 and chromosome Y, respectively. The cells cultured in SM showed a deletion of chromosome 8 and chtb(X)(q21) in one of 20 analyzed cells, which was likely a random loss of chromosomes due to technical preparation of the slide. The sample 7 had an addition of chromosome 2 in half of the PS-expanded cells in passage 7, while the other cultures derived from the same sample showed a normal karyotype. In four of the seven analyzed samples, there were too few metaphases in the PS culture in passage 7. It was correlated with their declined proliferation compared to the SM cultured counterparts as shown in Fig. 3B. Interestingly, mutations occurred in both samples only in passage 7 but not in passage 3 suggesting that these chromosomal abnormalities were induced during the ex vivo expansion.

Discussion

The xeno-free and serum-free medium PS has been used to manufacture ADMSCs and umbilical cord-derived MSCs in some clinical trials [34, 40, 41]. However, there is only limited data reporting the immunophenotype of MSCs in this medium so far. Recently, our group observed that ADMSCs cultured in PS exhibited evaluated levels of negative markers [35]. For the first time, we demonstrated that PS induced HLA-DR upregulation in these cells. It is a concern regarding the minimal criteria for MSCs established by the ISCT [27, 28]. An increased HLA-DR expression was also observed in GMP manufactured bone marrow-derived MSCs (BMMSCs) using either human sera or platelet lysate supplemented media [42]. The authors reported that the HLA-DR expression was unpredictable and dynamic over time in those cases [43]. Our data suggested that donor age was correlated with the increased signal of HLA-DR. In line with this observation, PS only induced its overexpression in adult MSCs, but not in those derived from prenatal tissues such as the umbilical cord [35].

Like PS, SM was able to support the isolation and expansion of ADMSCs [35]. Therefore, we performed a comparative study of these two xeno-free and serum-free culture media using the same samples and a common protocol. Our data demonstrated that both media were capable to differentiate ADMSCs into adipogenic, osteogenic, and chondrogenic lineages. However, SM supported cell growth and prevented ADMSCs from senescence better than PS. Interestingly, only PS medium triggered the expression of HLA-DR and displayed lower expression of CD90 and CD105 in passage 7, while SM medium showed no changes in marker profile during culture time. This indicates that PS itself should be the reason for the observed changes in the cells. Indeed, several factors in culture media can directly influence the MSC phenotype. The expression of HLA-DR can be induced in vitro by incubation of MSCs with IFN-γ or other pro-inflammatory cytokines [42–44] and basic fibroblast growth factor (bFGF) [45, 46]. In addition, CD105 level reduced in BMMSCs after pro-longed culture under a serum-free condition or during chondrogenic differentiation in the presence of transforming growth factor beta 3 [47, 48]. However, it remains elusive which factors in the PS medium caused the altered characteristics of ADMSCs.

The consequence of an HLA-DR upregulation on the bioactivity of MSCs remains controversial. Previous reports suggested that the HLA-DR overexpression on BMMSCs did not change their identity and potency [42, 43]. In contrast, our data indicated that in addition to HLA-DR induction, ADMSCs changed their karyotypes and earlier senescence upon a prolonged culture in the PS medium. For comparison, the SM medium showed a more stable karyotype and higher proliferation capacity under the same condition.

Furthermore, HLA-DR-expressing ADMSCs in the PS culture might increase the risk of alloimmune response as previously discussed for allogenic HLA-DR+ MSCs [49]. Several animal studies have shown that MSCs triggered immune reactions and resulted in a rejection of the administrated cells in HLA-mismatched mice [29, 50, 51], pigs [30], horses [31, 32], and monkeys [52]. On the other hand, INF-γ significantly enhanced the immunosuppressive effects of primed MSCs cells, although it induced an upregulation of HLA-ABC and HLA-DR [53–55]. Thus, the impact of HLA-DR expression on clinical use needs to be further investigated.

Conclusion

The xeno-free and serum-free PS medium was unsuitable for the expansion of ADMSCs, because it induced HLA-DR expression and led to decreased cell growth and proliferation as well as abnormal karyotypes in contrast to the SM medium. In addition, we provided evidence for a correlation between the increased expression of negative markers and the donor’s age in MSCs. These findings suggest the importance of culture media for the features of MSC-based products and emphasize the necessity of intensive quality control of these cells for a successful translation from bench to bedside.

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Authors’ Contributions Concept and design of the study: PTMD, VTH, XHN, NTL. Acquisition of data: PTMD, VTH, HTHB, LMH, LHT. Analysis and interpretation of data: PTMD, VTH, HTHB, LMH, HPN, LHT, HTTT, XHN. Drafting or revising the manuscript: PTMD, VTH, XHN, NTL, HTHB, LMH, HMD, HPN, LHT, HTTT. All authors have approved the final article.

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Data Availability The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Declarations

Ethics Approval and Consent to Participate Sample collection and data analysis were approved by the Ethics Committee of Vinmec Healthcare System and were carried out in accordance with the Declaration of Helsinki. Informed consent was signed by all donors before any study-related interventions took place.

Consent for Publication Not applicable

Competing interests The authors of the study are employed by the not-for-profit Vinmec Healthcare System.

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