Stromal cell therapy

A robust and standardized method to isolate and expand mesenchymal stromal cells from human umbilical cord

Pia Todtenhaupt1,2, Laura A. Franken1, Sophie G. Groene1,2, Marcella van Hoolwerff2, Lotte E. van der Meeren3,4, Jeanine M.M. van Klink2, Arno A.W. Roest5, Christiaan de Bruin6, Yolande F.M. Ramos1, Monique C. Haak7, Enrico Lopriore2, Bastiaan T. Heijmans1, Melissa van Pel8,9,*,**

1 Molecular Epidemiology, Department of Biomedical Data Sciences, Leiden University Medical Center, Leiden, The Netherlands
2 Neonatology, Willem-Alexander Children’s Hospital, Department of Pediatrics, Leiden University Medical Center, Leiden, The Netherlands
3 Department of Pathology, Leiden University Medical Center, Leiden, The Netherlands
4 Department of Pathology, Erasmus Medical Center, Leiden, The Netherlands
5 Pediatric Cardiology, Willem-Alexander Children’s Hospital, Department of Pediatrics, Leiden University Medical Center, Leiden, The Netherlands
6 Pediatric Endocrinology, Willem-Alexander Children’s Hospital, Department of Pediatrics, Leiden University Medical Center, Leiden, The Netherlands
7 Fetal Medicine, Department of Obstetrics, Leiden University Medical Center, Leiden, The Netherlands
8 NecestGen, Leiden, The Netherlands
9 Department of Internal Medicine, Leiden University Medical Center, Leiden, The Netherlands

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ABSTRACT

Background aims: Human umbilical cord-derived mesenchymal stromal cells (hUC-MSCs) are increasingly used in research and therapy. To obtain hUC-MSCs, a diversity of isolation and expansion methods are applied. Here, we report on a robust and standardized method for hUC-MSC isolation and expansion.

Methods: Using 90 hUC donors, we compared and optimized critical variables during each phase of the multistep procedure involving UC collection, processing, MSC isolation, expansion and characterization. Furthermore, we assessed the effect of donor-to-donor variability regarding UC morphology and donor attributes on hUC-MSC characteristics.

Results: We demonstrated robustness of our method across 90 UC donors at each step of the procedure. With our method, UCs can be collected up to 6 h after birth, and UC-processing can be initiated up to 48 h after collection without impacting on hUC-MSC characteristics. The removal of blood vessels before explant cultures improved hUC-MSC purity. Expansion in Minimum essential medium supplemented with human platelet lysate increased reproducibility of the expansion rate and MSC characteristics as compared with Dulbecco’s Modified Eagle’s Medium supplemented with fetal bovine serum. The isolated hUC-MSC showed a purity of ~98.9%, a viability of >97% and a high proliferative capacity. Trilineage differentiation capacity of hUC-MSCs was reduced as compared with bone marrow-derived MSCs. Functional assays indicated that the hUC-MSCs were able to inhibit T-cell proliferation demonstrating their immune-modulatory capacity.

Conclusions: We present a robust and standardized method to isolate and expand hUC-MSCs, minimizing technical variability and thereby lay a foundation to advance reliability and comparability of results obtained from different donors and different studies.

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Introduction

Isolation and expansion of primary cells from tissues often results in heterogeneous cell populations that may differ widely per culture due to the absence of standardized protocols and large donor-to-donor differences. To allow comparability between cells obtained from different donors within and between study cohorts and to
generate reproducible results, standardized methods for cell isolation and expansion are key. Human umbilical cord-derived mesenchymal stromal cells (hUC-MSCs) are a cell type that is increasingly used as a therapeutic agent and research model [1–5]. Standardized methods for hUC-MSC isolation and expansion are currently lacking, impacting on comparability and reproducibility of results obtained in study cohorts.

MSCs are multipotent stromal precursor cells originating from the embryonic mesoderm [6,7]. They can be isolated from a variety of different adult and birth-associated tissues, such as bone marrow, adipose tissue, umbilical cord (UC), placenta, amnion and chorion [2,8,9]. As the isolation from birth-associated tissues poses less ethical concerns due to the non-invasive collection procedures, their use as a source for MSC, especially UC, has increased [5,9–11]. Even though an active field of research currently centers on hUC-MSCs, thus far there is no consensus on the optimal method to isolate, expand and characterize hUC-MSCs. This hampers comparability and reproducibility of results obtained from studies using hUC-MSCs.

The isolation and expansion of hUC-MSCs is a multi-step process. After birth, the UC is collected and stored under conditions that maintain cord quality for subsequent MSC isolation. After transfer to a cell culture facility, the UC is inspected and prepared for culture. The UC tissue is then cultured to allow for migration of MSCs out of the UC pieces. Upon completing subsequent MSC expansion, the cells are cryopreserved for downstream applications, including full characterization. Variable protocol choices in each phase of the procedure may impact hUC-MSC characteristics, which is why optimization and standardization at every step of the process is essential.

Many different protocols describing the isolation and expansion of hUC-MSCs are used and published. These protocols vary greatly in tissue collection and preparation methods, culturing conditions and hUC-MSC characterization [8,12]. Differences in processing of the starting material can introduce persistent and unpredictable variation in the MSCs. In current published protocols, the UC collection procedure, UC storage conditions and maximum storage duration are often poorly defined [13]. Subsequent processing of the UC differs largely per isolation protocol. Some protocols advise to remove blood vessels from the cord before MSC isolation [14–17], whereas others culture the cord including veins and arteries [10,18,19]. In addition, the MSC-isolation procedures from UC differ widely, where the culture of larger explants [18,20], smaller “fine-piece” explants [11,16,17,19,21] as well as enzymatic digestion [10,11,16,17,22] are among the most commonly used isolation techniques. During isolation and expansion, different culture media and various types and concentrations of serum and other medium supplements are used [13,23]. Also, seeding densities vary between protocols, yet harvesting confluence remains similar among protocols [19,23]. Variability at each of these steps can affect hUC-MSC characteristics, such as growth kinetics and morphology, among others, due to their effects on their molecular profile (e.g., transcriptome, methylene) [8,11,15]. In order to generate reproducible results and compare hUC-MSCs obtained from different donors and studies, it is key to maintain the underlying molecular profile while reducing variability resulting from environmental factors.

MSCs are characterized by their adherence to plastic, the expression of cell surface markers CD105, CD73, and CD90, while lacking hematopoietic markers such as CD45, and their capacity to differentiate into adipocytes, osteoblasts and chondrocytes [24,25]. For hUC-MSC isolation, protocols generally have an adherence-based selection method [26]. Although, phenotype analysis and assessment of differentiation capacity varies greatly between studies, cell surface expression of CD73, CD90, CD105 and CD45 is consistently assessed. However, in-depth immunophenotypic analyses are highly variable and appear unmethodical. Besides, the used antibody concentrations and analysis strategies are often not standardized within or between research groups, thereby impairing comparison of the generated results [12,13,23,25–27]. Similarly, regarding the differentiation capacity of hUC-MSCs, a range of contradictory results have been published. Although some studies present an excellent differentiation capacity, others describe the inability to differentiate into any of the three lineages, leaving the actual capacity for trilineage differentiation indeterminate [19,21,28–30].

To enhance the robustness and reproducibility of hUC-MSC–derived results between donors within a study cohort and between studies from different research groups, it is crucial to standardize hUC-MSC isolation and expansion protocols. By comparing, selecting and optimizing variables at each step of this procedure, we have developed and standardized a method for the isolation, expansion and characterization of hUC-MSCs. Using an extended population of 90 donors, we isolated and expanded MSCs from UCs to show reproducibility of our method.

Materials and Methods

Ethical statement

hUCs were collected at the department of obstetrics at the Leiden University Medical Center in the Netherlands with ethical approval of the institutional medical ethical committee (P18.184). Written informed consent for the collection of hUC for research purposes was obtained from all parents. Mothers were included when diagnosed with a monochorionic twin pregnancy in the framework of the Twin-life study (International Clinical Trials Registry Platform ID NL7538) [31].

UC donor characteristics

All UC samples were derived from monochorionic twin pregnancies [31]. Per pregnancy, one individual of each twin pair was included in the analysis. The first presenting fetus of which the placental cord insertion during pregnancy was closest to the maternal cervix at first ultrasound was used for analysis, as this is considered a random characteristic. In total, UCs from 90 donors were included (female, 47 [52%]; delivery by cesarean section, 59 [66%]). The median gestational age at birth was 33.8 weeks (range, 25.9–37.0 weeks; interquartile range [IQR], 30.4–36.1 weeks). The median birth weight was 1853 g (range, 660–3330 g; IQR, 1376–2362 g).

UC collection and processing

At least 5 cm of the hUC was collected as soon as possible, with a maximum of 6 h after vaginal or cesarean delivery in collection buffer (phosphate-buffered saline [PBS]) supplemented with 0.38 μg/mL polymyxin B-sulphate (Sigma-Aldrich, St. Louis, MO, USA), 20 μg/mL kanamycin (Gibco/Thermo Fisher Scientific, Waltham, MA, USA), 10 μg/mL penicillin/streptomycin (Lonza, Basel, Switzerland) and 1 μg/mL Amphotericin-B (Sigma-Aldrich). The hUC was kept in collection buffer at 4°C until processing. Before processing, characteristics of the hUC were documented, including appearance, length of the collected hUC (centimeters), thickness (centimeters), cooking (intensity scale 1–5), the amount of blood clots (intensity scale 1–5) and edematous Wharton’s Jelly (intensity scale 1–5). Thereafter, the hUC was cut into 2-cm segments and washed in sterile PBS until all superficial blood clots were removed. The hUC pieces were transferred onto a Petri dish on ice and cut longitudinally to expose the umbilical arteries and vein. The blood vessels and remaining blood was removed as much as possible. Additional superficial incisions were made on the inside of the cord to increase the tissue surface and adherence. Approximately six dissected hUC pieces of ~1 cm² surface with a height of 3–5 mm were transferred into a sterile 10-cm Petri dish with the inside of the cord (Wharton’s Jelly) facing down. Cord pieces were left to adhere to the plastic for 20 min at room temperature.
(RT) before ~8 mL of culture medium was added. The culture medium level did not exceed the height of the hUC pieces to prevent detachment of the cord from the dish. Subsequently, the dishes were incubated in a humidified atmosphere at 37°C with 5% CO₂.

**hUC-MSC isolation and culture**

Minimum essential medium α (αMEM) GlutaMAX (Gibco) was used as standard culture medium supplemented with 100 μg/mL penicillin/streptomycin (Gibco) and 5% PLTGOLD human platelet lysate (hiPL) (Mercck, Rahway, NJ, USA). Where stated otherwise, Dulbecco’s Modified Eagle’s Medium (DMEM) GlutaMAX (Gibco) supplemented with 100 μg/mL penicillin/streptomycin and 10% fetal bovine serum (FBS) (Sigma) was used. Culture medium was changed twice a week.

Outgrowth of the hUC-MSCs is mainly visible at the perimeter of the explants. When more than one-half of the forming hUC-MSC patches showed a confluence of >70%, the explants were removed, and the dish was washed twice with PBS. hUC-MSCs were dissociated using TrypLE Select (Gibco). Total cell count and viability were established using a hemocytometer. Subsequently, the cells were seeded into a culture flask at a density of 2500/cm². When the hUC-MSC reached a confluence of >70%, the hUC-MSCs were dissociated from the plate using TrypLE Select. Passage 1 cells were used for experiments unless stated otherwise. The absence of mycoplasma was confirmed by polymerase chain reaction (PCR).

**Phenotype assessment by flow cytometry**

hUC-MSCs were stained with saturating quantities of CD73 (AD2, BV421), CD90 (5E10, PerCP-CyTM5.5), CD105 (SN6, FITC), CD45 (2D1, APC-H7) and CD31 (WM59, APC) monoclonal antibodies and the LIVE/DEAD stain kit (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s protocol. Cells were analyzed using the FACSCanto I Flow Cytometer using FACS Diva software for acquisition. Flow cytometer settings were performed as previously described [32]. Data were analyzed using BD FlowJo, version 10, software, as shown in Supplementary Figure 1.

**Adipogenic differentiation and Oil-Red-O staining**

For adipogenic induction, 2500 cells/cm² (bone marrow [BM]-MSCs) or 640 cells/cm² (hUC-MSCs) were cultured in adipogenic induction medium containing DMEM GlutaMAX supplemented with 100 μg/mL penicillin/streptomycin, 10% FBS, 10 μg/mL dexamethasone (Sigma-Aldrich), 10 μg/mL insulin (Sigma-Aldrich), 5 μmol/L 3-sobutyl-1-methylxanthine (Sigma-Aldrich), 50 μmol/L indomethacin (Sigma-Aldrich) and 1 μmol/L rosiglitazone (Sigma-Aldrich). The medium was refreshed twice a week. After 21 days of differentiation, the lipids and neutral triglycerides were stained using Oil-Red-O. For this purpose, 2.5 × 10⁵ MSCs were pelleted in 15 mL of propylene conical tubes (260 xg, 5 min). After overnight incubation, the standard culture medium was replaced with a serum-free chondrogenic induction medium consisting of DMEM GlutaMAX, 100 μg/mL penicillin/streptomycin, 5% PLTGOLD human platelet lysate (Sigma-Aldrich), 100 μg/mL dexamethasone, 10 μg/mL insulin (Sigma-Aldrich), 5 μmol/L 3-sobutyl-1-methylxanthine (Sigma-Aldrich), 50 μmol/L indomethacin (Sigma-Aldrich) and 1 μmol/L rosiglitazone (Sigma-Aldrich). After, the cells were incubated with Alizarin Red S solution (Sigma-Aldrich) for 10 min at RT. For removal of the excess staining, the cells were washed three times with milliQ before imaging with the Olympus BX53.

**Chondrogenic differentiation and Alcian Blue staining**

Chondrogenesis was induced in 3D pellets as previously described [34]. For this purpose, 2.5 × 10⁵ MSCs were pelleted in 15 mL of propylene conical tubes (260 xg, 5 min). After overnight incubation, the standard culture medium was replaced with a serum-free chondrogenic induction medium containing DMEM GlutaMAX, 10 μg/mL penicillin/streptomycin, 10 μg/mL L-dexamethasone, 0.1 μg/mL L-proline (Sigma-Aldrich), 100 μg/mL 3-sobutyl-1-methylxanthine (Sigma-Aldrich), 10 μg/mL ITS-Plus (Sigma-Aldrich) and 10 ng/mL hTGF-β1 (Tebu-bio, Ill de France, France) [34]. The chondrogenic culture medium was refreshed twice a week for a period of 5 weeks, of which from the second week onwards hypoxic conditions were applied. Thereafter, the diameter and surface area of the pellets was assessed microscopically using the Olympus BX53. The pellets were then fixed in 4% formaldehyde at 4°C overnight. Subsequently, the pellets were washed with 50% ethanol and stored in 70% ethanol until dehydration with an automated tissue processor and paraffin embedding. Sections of 5 μm were mounted on a glass slide and deparaffinized by incubating the slides twice in Histo-Clear (10 min; National Diagnostics, Atlanta, GA, USA), twice in 100% ethanol (5 min), once in 96% ethanol (3 min), once in 70% ethanol (3 min), once in 50% ethanol (3 min) and once in milliQ (3 min). To visualize the deposited glycosaminoglycans, the slides were primed for 5 min in 0.1 N HCl (J.T. Baker, pH 1.5, 5 min), followed by a 30-min incubation in 1 A% Alcian Blue 8-GX solution in 0.1 N HCl (Sigma-Aldrich). Excess staining was removed by 10 min of incubation with 0.1 N HCl and rinsing followed by a 30-min incubation with milliQ. Counterstaining was performed with a subsequent incubation in Nuclear Fast Red (Sigma-Aldrich). After 5 min, the slides were rinsed with milliQ (5 min) and dehydrated with 50% ethanol, 70% ethanol, 100% ethanol and Histo-Clear (1.5 min each), before the slides were mounted with Pertex (HistoLab, Brea, CA, USA) and imaged with the Olympus BX53.

**RNA isolation, reverse transcription and real-time quantitative PCR**

Cells derived from monolayer cultures were harvested, washed and snap frozen as a pellet in liquid nitrogen. RNA was isolated using the Zymo Quick-RNA MicroPrep Kit (Zymo Research, Irvine, CA, USA) according to the manufacturer’s protocol. RNA concentration was assessed using Qubit RNA BR Assay Kits (Life Technologies, Carlsbad, CA, USA). To synthesize cDNA, the Transcription First Strand cDNA synthesis kit (Roche, Basel, Switzerland) was applied according to the...
provided protocol using 200 ng of RNA as input. Each 10-μL PCR contained 2.5 μL Taqman Fast Advanced Master Mix (Applied Biosystems, Waltham, MA, USA), 0.5 μL of Taqman Gene Expression assay, 0.5 μL of CDNA and 6.5 μL of milliQ. Taqman Gene Expression Assay IDs were Hs01047975_m1 (RUNX2), Hs00939627_m1 (GUSB, Housekeeping gene), Hs01122454_m1 (YWHAZ, Housekeeping gene). The PCR was run on the QuantStudio 6 Flex (Applied Biosystems) for 2 min at 50°C, for 20 s at 95°C, followed by 40 cycles of 1 s at 95°C and 20 s at 60°C. Each condition was run in triplicate. The average of the technical replicates was only used for analysis when the standard deviation of the technical replicates did not exceed 0.5. The fold change was calculated by the 2^ [-ΔΔCT] method.

Inhibition of T-cell proliferation

Peripheral blood mononuclear cells (PBMCs) were isolated fromuffy coat of anonymous blood bank donors (Sanquin, Amsterdam, The Netherlands), using Leucosept centrifuge tubes according to the manufacturer’s protocol (Greiner Bio-One, Kremsmunster, Austria). Subsequently, the PBMCs were stimulated with Human T-Activator CD3/CD28 Dynabeads (Thermo Fisher Scientific) (5 μL/10^6 PBMCs) and cultured in the presence of 2 hUC-MSCs for 5 days at 37°C and 5% CO₂. PBMC to hUC-MSC co-culture ratios of 1:2, 1:8, 1:32, 1:128 and 1:512 were assessed. Unstimulated PBMCs served as a negative control, whereas CD3/CD28 stimulated PBMCs served as a positive control. Thereafter, the cultures were incubated overnight with 3H-thymidine. Incorporation of the 3H-thymidine was measured and displayed as a percentage of the positive control.

Data analysis and visualization

The associations between MSC characteristics and the attributes of donors and hUCs were quantified using the Spearman’s rank correlation. The P-values were Bonferroni corrected and considered statistically significant when < 0.05. Differences in population doubling time, population doubling level, surface marker expression, cumulative cell count and viability between culture conditions (with/without blood vessels; αMEM-hPL/DMEM-FBS) were tested using a two-sided paired t-test. Differences in surface marker expression and T-cell proliferation inhibition capacity between short- and long-term cryopreserved hUC-MSCs were tested using a two-sided unpaired t-test. Statistical analyses were performed using R software, version 4.1.0. Graphs and figures were created using R Software, version 4.1.0, and BioRender (https://biorender.com).

**Results**

**Standardized isolation and expansion of MSC from 90 UCs**

To develop a method for the standardized isolation and expansion of hUC-MSCs, we used a series of cords obtained from 90 donors. The isolation and expansion of MSCs from UC is a multi-step process spanning multiple days. Herein, we summarize the isolation and expansion method after protocol optimization (Figure 1).

UCs obtained from 90 donors after cesarean or vaginal delivery were collected within 6 h (median 0.6 hours) after birth and stored in an antibiotic/antimycotic solution (collection buffer). The MSC isolation process was initiated within 48 h (median, 17 h) after collection (Table 1). A morphological assessment of the UC was performed. Thereafter, the blood vessels and residual blood were removed and UC pieces of ~1 cm² (UC explants) were plated onto a culture dish.

After approximately 3 days of culture in the presence of human platelet lysate supplemented culture medium, it was observed that the first MSCs migrated out of the UC explants. In the subsequent week, the number of hUC-MSCs surrounding the cultured explants increased and confluent colonies were formed. Blood cells were observed throughout the first week of culture and diminished upon subsequent culture medium changes (Figure 2).

After a median culture period of 10 days (IQR, 9.0–11.0 days, Table 1), more than 50% of the MSC colonies reached a confluence of >70%. At this stage, UC pieces were removed and passage 0 (P0) hUC-MSCs were transferred to a culture flask and expanded for an additional passage. The number of cells obtained per UC explant (median 21.5 × 10^3 cells) varies greatly between the individual donors despite the highly comparable initial amount of tissue used for culture initiation (IQR, 14.3–39.7 × 10^3 cells, Table 1). The viability of the P0 hUC-MSCs was comparable between the donors (median, 97.2%; IQR, 95.9–98.4%, Table 1).

The passage 1 (P1) cultures were assessed daily for confluence. When reaching a confluence of >70% (median, 4 days; IQR, 4–4 days), hUC-MSCs were harvested. The population doubling level (PDL; median, 4.8 doublings; IQR, 4.5–5.1 doublings) and population doubling time (PDT; median, 0.9 days; IQR, 0.8–0.9 days) were comparable across the 90 hUC-MSC isolates (Table 1). A median of 99% of the cells (IQR, 98.6–99.2) expressed the cell surface markers CD73, CD90 and CD105 and did not express CD45 and CD31 (Table 1). The cell surface marker expression did not differ (P = 0.59) between short- (mean, 0.6 months) and long-term (mean, 21 months) cryopreserved samples (Supplementary Figure 2A). Combined, this indicates that this method allows for the reproducible isolation and expansion of hUC-MSC across UC donors.
MSC characteristics are stable and yield robust results across all donors. To assess the potential impact of donor characteristics on the hUC-MSC cultured using our method, birth weight, gestational age, sex and mode of delivery were correlated with outgrowth parameters and hUC-MSC properties. No associations were found between the donor characteristics and the number of hUC-MSCs obtained per UC explant or the hUC-MSC viability, indicating the robustness of our protocol. Notably, a lower gestational age \((P = 2.6 \times 10^{-4})\), a lower birth weight \((P = 2.1 \times 10^{-3})\) and delivery by cesarean section \((P = 8.7 \times 10^{-5})\) significantly correlated with decreased number of days needed to conclude hUC-MSC outgrowth in P0 (Figure 3).

**Table 1**

<table>
<thead>
<tr>
<th>Umbilical cord collection and MSC characteristics.</th>
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<td>Characteristic</td>
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<tr>
<td>Umbilical cord collection</td>
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<tr>
<td>Time between birth and umbilical cord collection, h</td>
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<tr>
<td>Time between collection and umbilical cord processing, h</td>
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<tr>
<td>Mesenchymal stromal cells</td>
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<td>Days in passage 0</td>
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<td>Cells obtained per UC piece</td>
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<td>Passage 0 ((\times 10^3))</td>
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<td>Cumulative cell count per UC</td>
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<td>UC piece Passage 1 ((\times 10^5))</td>
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<td>Viability passage 1</td>
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<td>PDL passage 1</td>
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<td>CD73(^+) passage 1, %</td>
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<td>CD90(^+) passage 1, %</td>
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<td>CD105(^+) passage 1, %</td>
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<td>CD45(^+) passage 1, %</td>
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<td>CD31(^+) passage 1, %</td>
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<td>CD73(^+)/CD90(^+)/CD105(^+), %</td>
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</table>

Characteristics of the umbilical cord collection and of the mesenchymal stromal cells are shown.

MSC, mesenchymal stromal cell; PDL, population doubling level; PDT, population doubling time.

**hUC-MSCs culture characteristics show limited donor-to-donor variability**

Donor characteristics such as birth weight, gestational age, sex and mode of delivery could affect hUC-MSC culture characteristics and may thereby impede comparability of cells obtained from different donors. To assess the potential impact of donor characteristics on the hUC-MSCs cultured using our method, birth weight, gestational age, sex and mode of delivery were correlated with outgrowth parameters and hUC-MSC properties. No associations were found between the donor characteristics and the number of hUC-MSCs obtained per UC explant or the hUC-MSC viability, indicating the robustness of our protocol. Notably, a lower gestational age \((P = 2.6 \times 10^{-4})\), a lower birth weight \((P = 2.1 \times 10^{-3})\) and delivery by cesarean section \((P = 8.7 \times 10^{-5})\) significantly correlated with decreased number of days needed to conclude hUC-MSC outgrowth in P0 (Figure 3).

**Time limits for UC collection and storage**

Cord handling before MSC isolation is a two-step process. After birth, the UC is cut to discontinue the blood flow between placenta and newborn. Thereafter, the UC is cut from the placenta and placed in the collection buffer to decelerate dehydration and cell death. Second, the UC is stored at 4°C until processing. Since an increased collection and storage time may affect hUC-MSC characteristics such as viability, it is critical to establish time limits for these steps. We assessed whether a collection and storage time of up to 6 and 48 h, respectively, is associated with changes in hUC-MSC outgrowth efficiency and characteristics. Neither the time required for hUC-MSC outgrowth, retrieved number of cells per explant, viability nor the PDT, PDL or immunophenotype correlated with the time between birth, collection and processing (Figure 3). This indicates that it is feasible to collect the UC up to 6 h after birth and store the UC for up to 48 h after collection without altering hUC-MSC characteristics and outgrowth performance. It should be noted however, that 89% (80) of the UCs were collected within one hour after birth whereas merely 3% (3) of the UCs were collected 4 or more hours after birth, resulting in a lower sensitivity to test the effect of moderate delays in collection on hUC-MSCs culture.

**Variation in UC characteristics do not associate with hUC-MSC properties**

Every umbilical cord has a unique morphology (Figure 4). The extent of coiling, edematous Wharton’s jelly and blood vessel visibility vary greatly among UCs (Figure 4), and this may affect the full removal of the arteries and vein during UC dissection. Also, we observed that the presence of blood clots and high mucinousity is a predictive factor for blood cell contamination in the first week of culture. Differences in UC morphology and explant composition may introduce heterogeneity in culture conditions and as such affect hUC-MSC characteristics and purity. To assess the potential impact of UC composition on the hUC-MSCs obtained using our method, all UC characteristics were mapped before processing, including the extent of coiling, blood clots, mucinousity, edematous Wharton’s jelly and visibility of blood vessels. Next, these were correlated with the hUC-MSC characteristics, growth performance and population purity. No significant associations were identified. Neither the time required for hUC-MSC outgrowth, retrieved number of cells per explant, hUC-MSC viability nor the PDT, PDL or the expression of cell surface markers were affected by variations in UC morphology (Figure 3). Taken together, with our method hUC-MSC characteristics are stable and yield robust results across all hUC-types.

**Figure 2.** MSC outgrowth from the UC. Visualization of the outgrowth of MSCs from the UC. Explants are plated on a dish (10 cm) with the inside of the cord facing down. After approximately 3 days, individual MSCs become visible at the explant perimeter. Over time, the blood cell presence decreases and the MSCs expand, forming colonies along the explant perimeter. Before imaging, the cord was removed from the dish for optimal contrast. The location of the cord explant is indicated by gray lining. Solid arrows indicate MSCs. The boxes and dashed arrows indicate the location of which the enlarged image is depicted below. Scale bars indicate 200 μm.
random donors were each cultured with and without blood vessels. Subsequently, hUC-MSC characteristics and purity were compared between the three selected donors of each group. At P0, more blood cells were observed in the dishes with explants of which the blood vessels were not removed (Figure 5A). Also, an increased percentage of CD45⁺ leukocytes and CD31⁺ endothelial cells was observed in these cultures (Figure 5B). However, this percentage was highly variable between the three donors (90.6%–97.9%). Leukocytes and endothelial cells disappeared after P0 and were not detected in the P1 hUC-MSC populations (Figure 5B, Supplementary Figure 3C). hUC-MSCs derived from hUC explants with and without blood vessels exhibited the typical elongated, fibroblastic-like spindle-shaped morphology. The hUC-MSCs did not differ in size, granularity or complexity between the two conditions (Figure 5A, Supplementary Figure 3). Also, the PDL and PDT were comparable, and the cultures derived from both dissection methods yielded comparable numbers of cells with a similar viability for at least three passages.

Thus, retaining the blood vessels in the explant during isolation did not impact proliferation capacity, cell counts and viability but led to a greater frequency of leukocytes and endothelial cells at P0. Therefore, we included the dissection of blood vessels from the UC-tissue before culture initiation in our method to avoid leukocyte and endothelial cell contamination. Small blood vessel fragments that might remain in the explants after dissection did not appear to affect hUC-MSC characteristics and can therefore be disregarded.

Culture in αMEM-hPL yields hUC-MSCs with a consistent growth performance and characteristics compared with DMEM-FBS

Culture medium and supplement may impact on MSC growth performance and characteristics [10,35,36]. To determine which culture medium and supplement yields the most consistent hUC-MSC growth performance and characteristics, we compared the two most commonly used medium-supplement combinations: αMEM supplemented with hPL (αMEM-hPL) and DMEM supplemented with FBS (DMEM-FBS) [37]. hUC-MSCs obtained from three different random donors were each cultured with and without blood vessels. Subsequently, hUC-MSC characteristics and purity were compared as culture medium in our method.

The time required for P0, and population doubling was increased in hUC-MSCs cultured in DMEM-FBS as compared with those isolated and expanded in αMEM-hPL, whereas the number of cells retrieved per explant as well as the PDL were decreased (Figure 6CD). Consequently, explants cultured in DMEM-FBS yielded considerably less hUC-MSCs (Figure 6B). Moreover, slightly lower viability was observed in DMEM-FBS-hUC-MSCs until passage 3, at which time point the viability of the αMEM-hPL-hUC-MSCs also started to decline (Figure 6E).

An increased percentage of CD45⁺ leukocytes and CD31⁺ endothelial cells was observed in P0 of DMEM-FBS-hUC-MSCs as compared with P0 αMEM-hPL-hUC-MSCs (Figure 6F, Supplementary Figure 4). In conclusion, the isolation and expansion in αMEM-hPL resulted in hUC-MSCs with a homogenous morphology and yielded a greater number of cells, increased viability and lower PDT as compared with DMEM-FBS. Therefore, αMEM supplemented with hPL was selected as culture medium in our method.
Trilineage differentiation capacity of hUC-MSCs as compared with BM-MSCs

MSCs are generally characterized by the capacity to differentiate into adipocytes, osteoblasts, and chondrocytes (Figure 7A) [25]. Therefore, we assessed the differentiation capacity of hUC-MSCs obtained from three random donors. BM-MSCs were used as positive control and undifferentiated MSCs were used as negative controls.

Over the course of the 3 weeks of adipogenic differentiation, the morphology of hUC-MSCs changed from a typical fibroblastic-like morphology towards highly irregular shaped cells with notably granular cytoplasm. Also, a greater content of neutral triglycerides and lipids as compared with the undifferentiated control was detected with Oil-Red-O staining. However, the typical fat storage in vesicles was absent in differentiated UC-MSCs, whereas this was observed in the differentiated BM-MSCs (Figure 7B).

After 3 weeks of osteogenic differentiation, hUC-MSCs showed no calcium deposition and little ALP activity. In contrast, differentiated BM-MSCs show a considerable amount of calcium deposition and ALP activity (Figure 7B). The hUC-MSCs kept expanding throughout the differentiation process and did not show signs of growth inhibition, as normally seen in these kinds of cultures. Gene expression analysis of the early osteogenic transcription factor RUNX2 was, however, increased in all differentiated hUC-MSC lines as compared with the control, compatible with early activation of the differentiation pathway. The differentiated BM-MSCs did not show elevated gene expression for RUNX2, possibly because their differentiation stage had passed the phase of RUNX2 upregulation (Figure 7C).

Prolonged three-dimensional chondrogenic differentiation over a 5-week period resulted in hUC-MSCs featuring an increased deposition of matrix as compared with the undifferentiated control cells. This was visualized by the staining of the main matrix components of cartilage tissue, glycosaminoglycans (Figure 7B). Also, the size of the three-dimensional pellet was consistently increased for all differentiated pellets as compared with the control pellets, indicating an increased deposition of extracellular matrix. The amount of matrix deposition reached after differentiation of BM-MSCs however was considerably more prominent as compared with UC-MSCs.
Thus, UC-MSCs appear to initiate differentiation when exposed to differentiation factors, although did not effectively complete differentiation as compared with BM-MSCs in the given time using the currently available protocols for MSC differentiation.

Isolated hUC-MSCs suppress T-cell proliferation

To assess the capacity of the hUC-MSCs to inhibit T-cell proliferation, hUC-MSCs obtained from 10 donors were co-cultured with CD3/CD28-stimulated T-cells. Co-culture with hUC-MSCs suppressed T-cell proliferation in a dose-dependent manner. With a hUC-MSC:PBMC co-culture ratio of 1:2, 13% of the proliferative capacity was retained relative to the control (Figure 7E). The capacity to inhibit T-cell proliferation did not differ between long-term (mean, 34 month) cryopreserved hUC-MSCs and short-term (mean, 12 month) cryopreserved hUC-MSCs (Supplementary Figure 2C), demonstrating that the immune-modulatory capacity of the hUC-MSCs isolated and expanded using our standardized method is independent of cryostorage duration.

Discussion

We developed a robust and standardized method for the isolation and expansion of hUC-MSCs to enable the comparison of hUC-MSC characteristics of different donors between future studies. To this end, we have optimized and standardized each step in the process of
MSC isolation and expansion from UC. In addition, by investigating the impact of donor heterogeneity on hUC-MSC characteristics, we were able to confirm robustness of our developed method across all donors.

We first specified time limits for the collection and subsequent storage duration of the UC before processing. After birth, a collection time of up to 6 h with an additional storage time of up to 48 h does not affect hUC-MSC isolation efficiency or characteristics, including proliferation capacity, viability and phenotype. In previously published protocols, the duration between birth and UC collection is rarely recorded [12,38]. The storage duration is more frequently described [13] and varies between 2 h [39] and 48 h [40]. Applying the presented time limits ensures that collection and storage duration does not impact on proliferation capacity, viability, and phenotype of the hUC-MSCs.

Next, we optimized the processing and isolation procedure of MSCs from UCs. Based on previously published results, we preselected the "plate-and-wait" technique to isolate UC-MSCs. With this method, explants are cultured without previous cellular dissociation by enzymatic treatment. We chose this method since enzymatic digestion has been reported to induce proteolytic stress, can result in cellular damage, and decreases cell viability [13,15–18,41,42]. Additional limitations of enzymatic digestion include degradation of cell surface receptors, alteration of cellular function, and increased PDT indicative of cellular ageing [14,16,18]. Taken together, these data favored our selection for the "plate-and-wait" technique.

In addition to the isolation procedure, the UC-processing procedures vary greatly between protocols. Although some methods include the removal of blood vessels before explant culture, others dismiss blood vessel removal as further tissue manipulation is more labor-intensive and holds a greater risk of culture contamination [13,18,43]. We found an increased number of endothelial cells and leukocytes in the hUC-MSC isolates at P0 upon retaining blood vessels in the UC explants. In line with this observation, another study employing a protocol without blood vessel removal, described residual contamination of endothelial cells in 30% of the samples at P1 [18]. In our hands, blood vessel removal before MSC isolation resulted in...
in a reduction of contaminating cells, including endothelial cells and leukocytes, without adverse effects on hUC-MSC characteristics and is therefore included in our method.

Another aspect we addressed is the influence of medium and medium supplements, as these are fundamental factors of variability in MSC isolation and expansion and can affect various MSC characteristics. Different media and media supplements have been used to culture MSCs [11,29,43,44]. Thus far, FBS is predominantly used as basal medium supplement. However, more recent studies are increasingly using hPL [37]. We selected αMEM-hPL as culture medium because we observed that culture in αMEM-hPL yields hUC-MSCs with a consistent growth performance and consistent characteristics as compared with DMEM-FBS. Likewise, previous studies also described favorable cell characteristics including increased proliferative capacity, decreased PDT and higher viability when comparing hPL with FBS in cultures of hUC-MSCs, BM-MSCs and adipose-tissue derived MSCs [10,35,36]. Furthermore, FBS shows a high batch-to-batch variability, requiring extensive and time-consuming comparability testing when a new batch is selected [45]. In contrast, hPL allows for increased batch-to-batch consistency and eliminates the risk of introducing animal-originating pathogens [46,47]. Also, in the view of potential clinical application of our expansion method, the use of animal components should be excluded where possible. Taken together, hUC-MSC culture in αMEM-hPL increases reproducibility and thereby comparability of results.

In addition, we standardised cell density and confluence at harvest as these can affect functional characteristics of MSCs. Seeding densities of $<500$ cells/cm$^2$ result in contact deficiency, whereas cells plated at a high density ($>5000$ cells/cm$^2$) are contact inhibited after a short culture period and must be passaged more frequently. Seeding densities of hUC-MSCs vary greatly in currently published protocols, describing densities from 100 cells/cm$^2$ to up to $1 \times 10^6$ cells/cm$^2$ [21,22,24]. To avoid frequent enzymatic treatment and to avoid contact deficiency, plating densities of $2.3 \times 10^3$ cells/cm$^2$ are recommended in the literature [23]. Based on this, we selected a plating density of $2.5 \times 10^3$ cells/cm$^2$. In contrast, the confluence for passaging/harvesting is stably described in protocols to be around 80% [15,19,40,43]. Based on local protocols, we used a harvesting/passaging confluence of $\sim70\%$ to avoid contact inhibition of the highly proliferative hUC-MSCs and to obtain consistent growth results.

We used our standardized method to isolate and expand MSCs of 90 hUCs. The time from culture initiation to harvest at P0 was approximately 10 days and P1 population doubling showed a median of 21.6 hours. Others described a P0 duration of $\sim14$ days [19,21] and a PDT varying between 24.8 and 29.4 hours using hPL [19,21] and 40.7 h using human serum as culture medium supplement [44], making our method considerably faster. This may be due to the relatively high amount of premature born donors and standardization of our protocol. The isolated hUC-MSC have a viability of $>97\%$ during P0 as well as P1, which is comparable or greater as compared with other studies [19,43]. For immunophenotypic characterization, we applied a method using standardized instruments settings, fluorochrome compensation and antibody staining ensuring comparability of all results [32]. We detected a purity of $\sim98.9\%$ in P1 hUC-MSCs, which was consistent throughout all 90 hUC-MSC isolates and comparable or improved as compared with other studies [19,21,43], demonstrating reproducibility of our method.

Next to variable protocol and reagent selection, heterogenous donor characteristics, including sex, gestational age, mode of delivery and birth weight, introduce variability. Yet, we found that the variability in donor characteristics did not impact the characteristics of our hUC-MSC isolates. This is in line with a previous study that could not detect a difference in hUC-MSC isolation and characteristics between male and female donors [43], confirming the robustness of our culture method across donors. However, we found that a lower gestational age, a lower birth weight and the delivery by cesarean section were correlated with a decreased time to P0 harvest. This contrasts with a previous study reporting a decreased MSC isolation efficiency in MSCs derived from cesarean delivery as compared with vaginal birth [43]. We hypothesize that in our study the time to complete P0 is decreased, due to an increased migratory capacity of the MSCs obtained from donors born at an early gestational age. The correlations with mode of delivery and birthweight may be a secondary effect, deriving from the fact that pre-term neonates are lighter and more frequently born by Cesarean section.

UCs from different donors vary in features such as the amount of coiling and edematous Wharton’s jelly, which can impact UC dissection and explants composition [48,49]. To our knowledge, we are the first to investigate the effect of various UC morphologies on the MSC isolation parameters, characteristics and population purity. We found no correlation between different UC phenotypes and characteristics of the hUC-MSC isolates, ensuring the robustness of our protocol across donors.

After demonstrating robustness of our method across donor and UC-types, we aimed to functionally characterize the generated hUC-MSC isolates. To that end, we differentiated MSCs obtained from three random UC donors towards adipocytes, osteoblasts and chondrocytes. We found a reduced capacity of hUC-MSCs to differentiate toward each of the lineages as compared with BM-MSCs, which differentiate efficiently into all three lineages. Previous studies describing the differentiation capacity of hUC-MSCs show mixed results. Although some describe hUC-MSCs to fail differentiation towards adipocytes, osteoblasts and chondrocytes [28], others report that UC-MSCs have the capacity to differentiate into all three lineages [44,50]. Yet, studies comparing hUC-MSC and BM-MSCs invariably report a reduced differentiation capacity of hUC-MSCs [21,29,30]. hUC-MSCs have been described to have greater expression levels of pluripotency markers such as Oct-3/4, Sox-2 and SSEA-4 as compared with BM-MSCs, indicating their less-mature character [12,51]. Therefore, it is possible that hUC-MSCs initiate differentiation when exposed to differentiation factors but do not readily proceed to full differentiation. In that case hUC-MSC may require tailored protocols to ensure a stable, reproducible, and complete differentiation towards adipocytes, osteoblasts and chondrocytes.

The ability to suppress T-cell proliferation makes hUC-MSCs a promising therapeutic agent and subject of numerous clinical trials [52]. By performing T-cell proliferation inhibition assays, we were able to demonstrate this immune-modulatory capacity of the hUC-MSCs. Therefore, our standardized method may also contribute to the standardization of hUC-MSC manufacturing methods.

**Conclusions**

We developed a standardized and robust method for the isolation and culture of hUC-MSCs. This protocol was shown to be stable across 90 hUC donors and HUC morphologies. After birth, a collection time of 6 h with an additional 48 h in process can be applied without impacting hUC-MSC characteristics. The removal of blood vessels before explant culture improves hUC-MSC purity of the hUC isolates. The culture in αMEM-hPL increases reproducibility, expansion rate and improves MSC characteristics as compared with DMEM-FBS. Our established protocol yields hUC-MSCs with a high purity, proliferative capacity, and viability. With this method, we have set the next step in unifying protocol choices regarding the isolation and culture of hUC-MSCs and thereby lay a foundation to advance the reliability and comparability of results obtained from hUC-MSCs of different donors.

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Author Contributions

Conception and design of the study: MyP, BTH, PT, JMMK, AAWR, CDb, MCH, EL. Acquisition of data: PT, LAF, SGG, MVH. Analysis of the data: PT. Interpretation of data: PT, MyP, BTH, YFMR, LEM. All authors have contributed to drafting and revising the manuscript and approved the final article.

Declaration of Competing Interest

The authors have no commercial, proprietary or financial interest in the products or companies described in this article.

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Supplementary materials

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