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Inefficiency in macromolecular transport of SCS-based microcapsules affects viability of primary human mesenchymal stem cells but not of immortalized cells

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ABSTRACT Microcapsules made of sodium cellulose sulphate (SCS) and poly-diallyl-dimethyl-ammonium chloride (pDADMAC) have been employed to encapsulate a wide range of established cell lines for several applications. However, little is known about the encapsulation of primary cells including human mesenchymal stem cells (hMSCs). Human MSCs are of interest in regenerative medicine applications due to pro-angiogenic, anti-inflammatory and immunomodulatory properties, which result from paracrine effects of this cell type. In the present work we have encapsulated primary hMSCs and hMSC-TERT immortalized cells and compared their behavior and in vitro angiogenic potential. We found that, although both cell types were able to secret angiogenic factors such as VEGF, there was a marked reduction of primary hMSC viability compared to hMSC-TERT cells when cultured in these microcapsules. Moreover, this applied to other primary cell cultures such as primary human fibroblasts but not to other cell lines such as human embryonic kidney 293 (HEK293) cells. We found that the microcapsule membrane had a molecular weight cut-off below a critical size, which caused impairment in the diffusion of essential nutrients and had a more detrimental effect on the viability of primary cell cultures compared to cell lines and immortalized cells.

KEYWORDS Microencapsulation, immunoisolation, sodium cellulose sulphate, mesenchymal stem cells and therapeutic angiogenesis.
INTRODUCTION

Human mesenchymal stem (stromal) cells (hMSCs) are a rare non-hematopoietic cell population present in bone marrow and other tissues capable of self-renewal and multi-lineage differentiation in vitro. They have the ability to form a variety of connective tissues including bone, fat and cartilage and they demonstrate specific surface antigen expression \(^{(1),(2)}\). Nevertheless, their therapeutic benefit is mostly related to the secretion of a wide range of paracrine factors, which may have anti-inflammatory, immunomodulatory and pro-angiogenic effects and may contribute to stimulation of local progenitor cells and tissue regeneration \(^{(3)-(5)}\). These properties may offer a clinical benefit for the treatment of inflammatory and ischemic conditions such as inflammatory bowel disease, coronary artery disease and peripheral arterial disease (PAD) \(^{(6)}\). Although results from current stem cell therapies using MSCs are encouraging, important limitations remain and major problems include the poor rate of engraftment and limited cell survival after transplantation. One of the reasons could be the susceptibility of MSCs (specifically allogeneic cells) to immune-mediated cell destruction, mainly through antibodies and complement proteins \(^{(7)-(10)}\).

The use of biomaterials to encapsulate cells may overcome these problems and may enhance the therapeutic efficacy of stem cell transplantation. A combination of stem cell therapy and microencapsulation technology has three main potential benefits; a) immunoisolation of enclosed cells, b) cell retention at the transplanted site and c) long-term delivery of therapeutic factors.

Microcapsule systems with high mechanical strength and that induce low cytotoxicity and minimal inflammation, are advantageous. In this regard, microcapsules made of sodium cellulose sulphate (SCS) and poly-diallyl-dimethyl-ammonium chloride (pDADMAC) have been shown to be very stable and biocompatible \(^{(11)-(13)}\). Moreover, safety and proof of principle with this
approach have been demonstrated in a phase I/II clinical trial in patients with inoperable pancreatic cancer \cite{14,15}. Several established cell lines including epithelial cells \cite{16,17}, fibroblasts \cite{18}, hybridoma cells \cite{13,19} islet cells \cite{20,21} and neural stem cells \cite{22} have been successfully encapsulated using this system. However, little is known about the behavior of primary cells such as hMSCs encapsulated using this technology.

SCS-based microcapsules containing hMSCs may represent a system with potential for enhancing persistence and efficacy in a broad range of conditions. One example would be in the treatment of ischemic disorders by induction of therapeutic angiogenesis. In the present work, we have characterized the morphology and chemical structure of SCS-pDADMAC microcapsules, the cytotoxicity of the microcapsule components and the diffusion properties of its membrane. Moreover, we have encapsulated primary hMSCs and hMSC-TERT immortalized cells \cite{23,24} and studied their behavior and the \textit{in vitro} angiogenic potential over a period of 14 days in culture. In this study, we have also encapsulated primary human fibroblasts and the cell line human embryonic kidney 293 (HEK293) to provide further data about the behavior of primary versus cell lines.

\textbf{MATERIALS AND METHODS}

\textbf{Materials.} SCS (batch E12/1NEG2S13) with a sulphate-degree of substitution (ds) of 0.40 was purchased from Fraunhofer IAP (Potsdam, Germany) and Poly- (diallyl-dimethyl-ammonium chloride) or pDADMAC of low molecular weight 100-200 kDa (Lot# MKBK3094V) and ultra-low molecular weight <100 kDa (Lot# MKMF5290V) was obtained from Sigma-Aldrich.

\textbf{Cell culture.} Primary hMSCs were isolated from human bone marrow (Lonza, Walkersville, MD) and characterized based on the expression of CD105, CD73 and CD90 and lack of
expression of CD45, CD34, CD3, CD19, CD14 and HLA-DR and the tri-lineage differentiation capacity as stipulated by the International Society of Cellular Therapy (2) (Supporting Information Figure S1). A total of three different donors were used (n=3). These cells were cultured in standard complete medium consisting of α-MEM supplemented with 10% fetal bovine serum (FBS; Hyclone™, Thermo Scientific), 1% penicillin/streptomycin (Gibco, Invitrogen). Primary human (foreskin) fibroblasts (hFibroblasts) were obtained from ATCC and cultured in standard complete medium. HEK293 were cultured in D-MEM (high glucose) supplemented with 10% FBS, 1% penicillin/streptomycin and 2 mM L-Glutamine. The human MSC-TERT (hMSC-TERT) cell line has been established by ectopic expression of the catalytic subunit of human telomerase as previously described (23),(24). These cells express all known markers of primary MSC and form ectopic bone and bone marrow organ when implanted in vivo. Human MSC-TERT cells were cultured with α-MEM supplemented with 10% FBS and 1% penicillin/streptomycin. Human Umbilical Vein Endothelial Cells (HUVECs) were obtained from Lonza and cultured in EBM-2 basal media (Lonza) containing EGM™-2 Bulletkit™ (serum and growth factors) (Lonza). All cells were maintained on a humidified incubator at 37ºC and 5% CO₂.

**Procedure to encapsulate cells.** For microcapsule preparation, 2% (w/v) SCS solution was prepared by dissolving 2 g SCS in 100 mL of sterile distilled water with 0.9% (w/v) of sodium chloride. This SCS solution was then filtered-sterilized (0.45 μm pore size). Cells were mixed with the 2% (w/v) SCS solution to give a final concentration of 5x10⁶ cells/mL. The SCS solution containing cells was passed through an encapsulator (Inotech encapsulator IE-50R, Switzerland) using a 200 μm nozzle. Droplets of equal size were formed and eventually fell into a well-stirred solution of sterile pDADMAC. The solution of pDADMAC was composed of
0.86% (w/v) of low molecular weight (100-200 kDa) and 0.06% (w/v) of ultra-low molecular weight (<100 kDa) species. The microcapsule shell was formed through polyelectrolyte complexation (PEC) of the anionic SCS with cationic pDADMAC polymer, resulting in the formation of stable liquid-core microcapsules containing cells, as previously described (11),(25). The reaction time was adjusted to 4 min, and after that, microcapsules were washed 3 times with D-PBS (Gibco, Invitrogen) and 3 times with standard complete medium. All procedures were performed under sterile conditions. Microcapsules containing cells were stored in standard complete medium at 37°C and 5% CO₂. Medium was changed every 2-3 days. The same conditions were used to generate microcapsules without cells.

Cell proliferation assay. For cell proliferation studies, primary hMSCs and hMSC-TERT cells were seeded at a density of 24,000 cells/well in a 12-well plate and cultured with complete medium, serum-free medium, 50K-depleted medium and 50K-concentrated medium for the culture period of 9 days. 50K-depleted medium was obtained by filtration of complete medium (α-MEM containing 10%FBS) with an ultra centrifugal filter device of 50K cut-off (Amicon® Ultracel-50K, Millipore) and 50K-concentrated media was obtained from the retentate volume. Both media were collected and used for culturing of cells. Protein content of media was analyzed by 15% SDS-PAGE and Silver Staining (Fermentas Life Science).

Characterization of microcapsules

Fourier transform infrared (FTIR) spectroscopy. The chemical structure of SCS, pDADMAC and polyelectrolyte complex (PEC) was analyzed by infrared spectroscopy. Analysis was performed on crude SCS (solid form) and pDADMAC (20% (v/v) solution, Sigma-Aldrich). For PEC membrane formation, a solution of 1% (w/v) pDADMAC was added to 2% (w/v) SCS until polymerization occurred and an insoluble compound was formed. The spectra were acquired on
an FTIR-8300 spectrophotometer with an average of 20 scans, a resolution of 4 cm\(^{-1}\) and a wavenumber range of 600-4000 cm\(^{-1}\) at ambient temperature.

**Microscopic analysis of microcapsules.** The diameter of 50 microcapsules from three independent encapsulation procedures was measured using confocal laser scanning microscopy (CLSM; Zeiss LSM 510 Axiovert Inverted Confocal Microscope) to determine microcapsule size distribution and reproducibility. Scanning electron microscopy (SEM) was employed to study the ultrastructure of microcapsules. For visualization of the microcapsules’ outer surface and membrane cross-section, microcapsules were dehydrated through graded ethanol, sputter-coated with colloidal gold and observed with Hitachi S2600N Variable Pressure Scanning Electron Microscope. For images created using Back Scattered Electron Mode (BES) – the Back Scattered Electron detector was inserted into the system. In this case, microcapsules did not undergo a dehydration process and instead, they were directly visualized. The beam size was increased to at least 80, a higher kV was also used (20-25kV) and pressure was set at between 50-80 Pascals (Pa).

**Cytoperformance of encapsulated cells**

**Determination of viability and metabolic activity.** Cell viability on days 0, 1, 3, 7 and 14 days post-encapsulation was assessed using the LIVE/DEAD® viability/cytotoxicity assay (Molecular probes, Invitrogen) as recommended by the manufacturer. Microcapsules containing cells were visualized under CLSM (Zeiss LSM 510 Axiovert Inverted Confocal Microscope) using the excitation/emission wavelengths of ~528nm/~617nm (red fluorescence) and ~495 nm/~515 nm (green fluorescence). Scans throughout the whole microcapsule were taken using a step size of 5\(\mu\)m. These scans were then projected onto 2D-plane (z projection) using the software provided. As a control, encapsulated hMSCs were killed using pure ethanol, stained with LIVE/DEAD®
dyes and observed using CLSM. The metabolic activity of encapsulated cells immediately after
encapsulation and on days 3, 7 and 14 post-encapsulation, was assessed using AlamarBlue®
assay. AlamarBlue® (10% of the volume of the well) was added to each well containing 30 ± 3
microcapsules (5 experimental replicas in 96-well plates) and incubated for 21 h at 37°C
protected from the light. After incubation, 2 aliquots of 100μl per well were transferred to a 96-
well plate. Absorbance was measured at 550nm with wavelength correction at 595nm using a
microplate reader (Victor3 1420 multilabel plate counter). Results were expressed as mean ± SD
from 3 independent encapsulation processes.

Cytoskeleton organization. Samples of microcapsules containing cells were collected at days 0,
3, 7 and 14, fixed in 4% paraformaldehyde in D-PBS for 20 min and permeabilized with 0.1%
TritonX-100 (Sigma-Aldrich) for 5 min. The nucleus of the encapsulated cells was stained with
0.25mg/mL of Propidium Iodide (PI) (Sigma-Aldrich) for 10 min. Cell cytoskeleton was stained
for F-actin with 50μg/mL Phalloidin-FITC (Sigma-Aldrich) for 45 min at room temperature
protected from light. Microcapsules were visualized using CLSM (Zeiss LSM 510 Axiovert
Inverted Confocal Microscope) using excitation and emission wavelengths of ~495 nm/~515 nm
(green fluorescence) and ~528nm/~617nm (red fluorescence), respectively.

Cell localization. Localization of cells inside the microcapsules was investigated using CLSM
(Zeiss LSM 510 Axiovert Inverted Confocal Microscope) and LIVE/DEAD® dyes as above.
Scans throughout the whole microcapsule were taken using a step size of 5μm and images were
projected to an orthographic projection.

In vitro cytotoxicity assays. The biological response of primary hMSCs to microcapsule
material was assessed as stipulated by the International Organization for Standardization, ISO
10993-5:2009(E). Briefly, 5,000 cells/well in 48-well plates were cultured for 48h in direct
contact with sterile empty microcapsules (direct contact) and with extracts of the microcapsule device through diffusion (indirect contact). In addition, cells were incubated with 2% and 4% (w/v) SCS solution re-suspended in complete medium at ratios of 1:3 and 2:3 (SCS:media). In the three experiments, cytotoxicity was assessed 48h later using an AlamarBlue® assay, as recommended by the manufacturer. Furthermore, primary hMSCs were cultured in suspension with a solution containing sterile empty microcapsules in serum-free conditions. The ability of primary hMSCs to spread on the outer microcapsule membrane was studied using CLSM at the time of cell seeding and 24h after, using transmitted laser light and LIVE/DEAD® reagents (z-projection and orthographic projection).

**Molecular weight cut-off of microcapsule membrane.** The molecular weight cut-off (MWCO) of the microcapsule membrane was determined by incubating 40-50 microcapsules in 1 mL of D-PBS, supplemented with 0.5 mg/mL of FITC-labeled dextran (10, 20 and 70 kDa; Sigma-Aldrich), 40 μg/mL of FITC-coupled chicken anti-rabbit IgG (or IgY, 180kDa; Abcam) and 0.5 mg/mL of FITC-coupled BSA (66.5 kDa, Sigma-Aldrich). After 20h, FITC-fluorescence inside and outside of the microcapsules was determined by CLSM (Zeiss LSM 510 Axiovert Inverted Confocal Microscope). The distribution of fluorescence intensity was obtained by performing a linear scan of microcapsules using the CLSM software.

**In vitro angiogenic potential of encapsulated cells.** Levels of Vascular Endothelial Growth Factor (VEGF) released from encapsulated cells were assessed by ELISA (R&D Systems) as recommended by the manufacturer. Seven days after encapsulation, microcapsules containing cells were re-suspended in fresh complete medium and samples of conditioned medium were collected at 24h, 48h and 72h for VEGF quantification. Similarly, VEGF levels were quantified from the conditioned medium of non-encapsulated cells (50,000 cells/well) cultured for 24h, 48h
and 72h. The ability of encapsulated cells to secrete factors that induce the formation of tube-like structures by HUVECs was investigated using a matrigel tube formation assay. After 7 days of culture, microcapsules containing cells were incubated with EBM medium (without addition of serum and growth factors) for 48h, after which time, the conditioned medium was generated and collected. Growth factor reduced matrigel basement membrane (BD Matrigel Matrix Growth Factor Reduced, BD Bioscience) was thawed on ice at 4°C overnight. A thin layer (110μL/well) was added to 48-well plates and allowed to solidify at 37°C. HUVECs (2.5x10^4 cells) were re-suspended with the appropriated media (3 experimental replicas: conditioned medium from encapsulated cells, EBM-2 medium for negative control and EGM-2 medium for positive control). Media containing HUVECs were added on top of the matrigel and incubated for 14h at 37°C. After incubation, four pictures per well were taken at random and a quantitative measurement of tube-like structures was carried out and normalized to the positive control. Tubules were considered as structures that were at least 4 times the length of a cell.

**Statistical analysis.** Statistical analysis of the results was performed using Graphpad Prism (Graphpad Prism 5 Software). The D’Agostino&Pearson test was used for testing the normal distribution of the samples. The results of multiple observations are represented as mean ± standard deviation (SD) (n=3). For two-sample data analysis, unpaired Student’s t-test was performed. For multivariate data analysis, group differences were assessed with one-way analysis-of-variance (ANOVA) with corresponding multiple comparison test, which are specified in the legend of each figure. Differences were considered statistically significant when p values were lower than 0.05.
RESULTS

Chemical and morphological characterization of microcapsules. Microcapsules made of SCS and pDADMAC are liquid-core microcapsules, which are formed when two oppositely charged polyelectrolytes (i.e. anionic SCS and cationic pDADMAC) interact in an aqueous solution. As a consequence, a solid-like precipitate called polyelectrolyte complex (PEC) is formed at the droplet surface, which acts as a semipermeable membrane\textsuperscript{(26)}. This semipermeable membrane may permit bidirectional diffusion of nutrients, oxygen and secreted factors but prevents the entry of bigger molecules (i.e. antibodies) and host immune cells (Fig. 1A). In the present work, we have characterized the morphology and structure of the SCS-pDADMAC microcapsules. Scanning electron microscopy (SEM) was used to study the overall microcapsule ultrastructure. Microcapsules made of SCS and pDADMAC are composed of a liquid-core of unreacted SCS polyelectrolyte enveloped by a semipermeable membrane made of PEC (Fig. 1B). SCS-pDADMAC microcapsules represent an osmotically interactive system\textsuperscript{(25)} and osmotic pressure differences may cause changes in capsule volume by movement of water. In figure 1B, microcapsules were visualized under SEM using the BSE mode to observe the natural structure of microcapsules. These samples were not dehydrated through graded ethanol and appeared as spherical structures with a smooth surface. It is proposed that the microcapsule’s liquid-core diffused out when they were subjected to the dehydration process during the preparation for SEM visualization, and as a result they appeared as collapsed spherical structures (Fig. 1C). In this image it is suggested that the dehydrated microcapsule is entirely formed by PEC membrane. FTIR spectrum was used to confirm the naïve structures of SCS and pDADMAC and the formation of the membrane by PEC after polymerization (Fig. 1D). The most characteristic peaks in the SCS spectrum are the bridge oxygen (C-O-C ether) and the C-OH stretching bands
observed as multiple peaks in the region of 990-1150 cm\(^{-1}\), and the strong bands of S=O and –SO\(_3\)\(^-\) at 1217 cm\(^{-1}\) and C-O-S at 808 cm\(^{-1}\). In the pDADMAC spectrum, a broad peak at 2100 cm\(^{-1}\), due to the –CN stretching, and a peak at 1473 cm\(^{-1}\) attributed to the –CH\(_3\) groups, were also identified. PEC membrane formation was confirmed by its solid-like structure and insoluble character compared to the unreacted liquid SCS and pDADMAC. The PEC spectrum includes peaks at 2100 cm\(^{-1}\) from –CN stretching and multiple peaks at the region of 990-1150 cm\(^{-1}\) from the C-O-C and C-OH groups.

Finally, microcapsule size distribution and reproducibility was studied by measuring the diameter of 50 microcapsules from 3 independent encapsulation processes. SCS-pDADMAC microcapsules produced by the vibrational technology are known to have a narrow size distribution \(^{(27)}\). Microcapsules produced with the conditions described in the materials and methods are highly reproducible and have a diameter of 556.99 ± 30.33 μm with a narrow size distribution (Fig. 1E).

**Survival of primary hMSCs is compromised after encapsulation.** Viability of encapsulated primary hMSCs was measured after encapsulation and on days 1, 3, 7 and 14 using LIVE/DEAD® viability/cytotoxicity assay. Primary hMSCs survived the encapsulation process, as they exhibited green fluorescence immediately after encapsulation (day 0). However, viability over time was compromised as was indicated by the appearance of red fluorescent cells (Fig. 2A). Metabolic activity measurements using AlamarBlue® assay confirmed these results (Fig. 2B). We found a pronounced reduction of metabolic activity of 64.44 ± 15.56% by day 3 compared to day 0, and this further decreased by day 7.

The localization of cells within the microcapsules was investigated using LIVE/DEAD® dyes and CLSM. Primary hMSCs remained in single cells and were distributed throughout the liquid-
core of the microcapsule immediately after encapsulation (Fig. 2C). However, we observed that
cells started to clump together a few hours after encapsulation, and they remained in clumps until
day 14. Furthermore, we observed that these cells failed to spread and localized close to the inner
microcapsule membrane (Fig. 2C and Video S1).

The morphology of hMSCs inside the SCS-based microcapsules was assessed by CLSM and
staining of F-actin and cell nuclei with Phalloidin-FITC (green) and PI (red), respectively (Fig.
2D). Primary hMSCs had a rounded morphology immediately after encapsulation and formed
clumps about 70-80 μm in diameter by day 3. Cells remained in a round shape within the
clumps, with poor cytoplasmic extensions around the cells (Fig. 2D). Transmitted laser light and
LIVE/DEAD® dyes were used to visualize the conformation of the cell clumps (Supporting
Information Fig. S2). Primary hMSCs clumps were initially formed by healthy cells, although
considerable cell death was detectable 3 days after encapsulation. At 14 days, these cell clumps
were mainly formed by a small core of viable cells (green) surrounded by dead cells (red).

**Microcapsule material is not cytotoxic to primary hMSCs.** The cytotoxicity of microcapsule
components to primary hMSCs was assessed 48 h after first contact with the material (by direct
or indirect contact), using an AlamarBlue® assay (Fig. 3A). Results showed that neither PEC,
which forms the microcapsule membrane (Fig. 3Ai), nor extracts released by microcapsules,
which may contain residual amount of polyelectrolyte (Fig. 3Aii), had a cytotoxic effect on
primary hMSCs. Moreover, metabolic activity of primary hMSCs was maintained in cells
cultured in direct contact with a solution of unreacted SCS (which forms the liquid-core of the
microcapsules) at concentrations of 2% and 4% (w/v) (Fig. 3Aiii). Interestingly, we found that
primary hMSCs were able to spread on the outer layer of the microcapsule membranes when
cultured in serum-free conditions for 24 h (Fig. 3B).
Immortalized hMSCs survive after microencapsulation. Results from LIVE/DEAD® assay showed that immortalized hMSC-TERT cells remained viable during the culture period tested (14 days), as most of the cells were stained with Calcein AM (Fig. 4A). Likewise, the AlamarBlue® assay showed that encapsulated hMSC-TERT cells maintained their metabolic activity during the 14 days (Fig. 4B). Human MSC-TERT cells also formed clumps, however, they were mainly composed of healthy cells stained with Calcein AM (green). These clumps were able to increase in size with little cell death (Supporting Information Fig. S2). Human MSC-TERT cells were found to move towards the inner microcapsule membrane similarly to primary hMSCs (Fig. 4C). Furthermore, hMSC-TERT cells were able to proliferate as seen by the higher cell density found inside of the microcapsule at day 14 compared to day 0 (Fig. 4A). Cells were able to spread on the inner membrane of the microcapsule as seen in orthographic projection in figure 4C and Video S2. Results from F-actin and cell nuclei staining showed that hMSC-TERT cells formed compact and well-organized clumps, which had different shapes (Fig. 4D). These cell clumps grew in size forming very dense cellular structures 14 days after encapsulation.

In order to provide further data on the behavior of primary cells and cell lines, we encapsulated other cell types including primary human fibroblasts and HEK293 cells (Supporting Information Fig. S3). Results from LIVE/DEAD® staining showed that a reduction in cell viability occurred in primary human fibroblasts but not in HEK293 cells.

Diffusion of macromolecules is impeded by the microcapsule membrane. The ultrastructure of the microcapsule membrane was determined by SEM. The microcapsule membrane appeared to be a very dense, smooth and compact structure, and large diffusion channels (i.e. pores) were not observed (Fig. 5A). The diffusion of 10, 20 and 70 kDa FITC-labeled dextrans, FITC-
coupled IgY (180kDa) and FITC-coupled BSA (66.5kDa) across the microcapsule membrane was investigated (Fig. 5B). Results showed that small molecules such as 10 kDa dextrans were able to permeate easily, while 20 kDa dextrans were able to permeate to an extent, as seen by a reduced fluorescence inside the microcapsules. Moreover, its diffusion did not improve at 96h (Supporting Information Fig. S4). Furthermore, BSA (66.5kDa), one of the most abundant globular proteins present in the FBS employed for the culture of cells, was not able to penetrate the microcapsule membrane and appeared adsorbed at the outer microcapsule membrane. Finally, large molecules including antibodies (i.e. 70kDa dextran and 180kDa IgY) were also not able to permeate the microcapsule membrane.

**Macromolecules present in complete medium have an impact in the growth of cells.**

Macromolecules present in complete medium were depleted based on size using a 50K-filter device. Protein content of media was analyzed by SDS-PAGE. Results showed that complete medium included proteins ranging from <1 to >180 kDa, while no protein content was detected by SDS-PAGE in the serum-free medium (Fig. 6A). We found that macromolecules larger than ~70kDa were retained in the concentrate while the concentration of proteins <50kDa were diminished as a consequence of the 50K-filtration step. This fractionation would mimic a possible delay and/or impediment on the entrance of macromolecules from the medium into the microcapsules.

Primary hMSCs and MSC-TERT cells were incubated with completed medium, serum-free medium and 50K-depleted medium, and cell proliferation was assessed 9 days after. Results showed that primary hMSCs failed to proliferate in the 50K-depleted medium and in serum-free conditions (Fig. 6B). Conversely, hMSC-TERT cells were able to proliferate in both serum-free conditions and 50K-depleted medium, although their growth was slower compared to the
complete medium (Fig. 6B). We found that hMSC-TERT cells cultured in 50K-depleted medium underwent ~9-fold increase in cell number compared to day 0. However, primary hMSCs showed a decrease in cell number when compared to day 0 (Fig. 6C).

**Encapsulated cells secrete angiogenic factors.** Primary hMSCs and hMSC-TERT cell line are known to secret angiogenic factors (31),(5),(28),(29). The release of functional angiogenic factors from the encapsulated cells was studied using a matrigel tube formation assay (30). We found that the conditioned medium (CM) from encapsulated primary hMSCs and hMSC-TERT cells was able to induce tube-like structures in HUVECs (Fig. 7A-B). Moreover, ELISA quantification of VEGF levels from non-encapsulated cells showed that both cell types were able to secrete this angiogenic protein. However, the release of VEGF by hMSC-TERT cells increased over time and at 72h (2374.42 ± 398.85 pg/mL) it was greater than primary hMSCs (1223.86 ± 684.56 pg/mL), probably due to the higher proliferation rate of this cell type (Supporting Information Fig. S5). Encapsulated primary hMSCs and hMSC-TERT cells also secreted VEGF (Fig. 7C), but again, hMSC-TERT cells performed better in terms of both quantity and duration of protein secreted.

**DISCUSSION**

Our results indicate the potential of SCS-pDADMAC microcapsules as devices for long-term delivery of angiogenic factors secreted by the encapsulated cells, while simultaneously protecting the cells from the contact with antibodies. This would be valuable for allogeneic stem cell transplantation in the field of therapeutic angiogenesis. However, we have found that the survival of primary cells is compromised after encapsulation.

To our knowledge, SCS-based microcapsule systems have been mainly employed for the encapsulation of established cell lines (13)-(22). However, the use of cell lines may be a concern for
the regulatory bodies when designing clinical trials, in particular, immortalized cells that are able
to form tumors \textit{in vivo}. For this reason, our interest focused on the encapsulation of primary
human cells such as MSCs, which are of interest in many clinical applications. In the present
work, we have also investigated the behavior of other primary cell types such as human
fibroblasts, and two cell lines including hMSC-TERT cells and HEK293 cells, for comparative
purposes.

Our results showed that the viability of primary cells such as hMSCs and hFibroblasts
encapsulated in this device was limited compared to the performance of established cell lines
such as hMSC-TERT and HEK293 cells. Likewise, hMSC had a reduced metabolic activity over
the 14 days of culture compared to hMSC-TERT cells. This observation suggests that primary
cells may find some elements of microcapsule environment detrimental whereas their
immortalized counterpart is able to survive in these conditions. Nevertheless, we found that
microcapsule material was not cytotoxic to primary hMSCs by either direct or indirect contact.

Intriguingly, we found that hMSC-TERT cells had extensive proliferation by day 14, and cells
occupied about 50\% of the microcapsule inner membrane. Uncontrolled proliferation from
immortalized cells could be worrisome from a safety point of view. Although these cells may be
protected from the host immune system by the microcapsule membrane, an uncontrolled
proliferation of these cells could cause a rupture of the microcapsule and release of the cells,
which could compromise the health of the patient. However, we did not observe an extensive
proliferation by the encapsulated primary hMSCs during the 14 days studied.

CLSM was used to study the localization of cells after encapsulation. These results showed
that primary hMSC were distributed as a single cell suspension throughout the microcapsule
immediately after encapsulation and remained with in rounded morphology for 14 days.
However, we observed a movement of cells out of the center towards the membrane, where nutrients and oxygen might be more readily available to cells, as previously reported (11). We suggest that this effect may be more likely to be caused by gravitational forces that pull the cells towards the bottom of the microcapsule, as the liquid-core of these microcapsules may not support the interaction or attachment of cells. Moreover, we observed that primary hMSC had a rounded morphology with no cytoplasmic elongations (which may be indicative of some kind of interaction with the material), suggesting that cells do not interact with the liquid-core of the microcapsules. In contrast, PEC that forms the microcapsule membrane, a solid-like precipitate (26), may provide better support for cell anchorage than the liquid-core of the microcapsules.

In this regard, it is important that the inner layer of the microcapsule membranes provides the correct adhesion signals to cells. Primary hMSCs are anchorage dependent cells, and as in many other cells types, they require integrin-mediated adhesion to extracellular matrix (ECM) proteins to display optimal metabolic functionality and viability. When this does not occur, cells may die through a mechanism called anoikis, or anchorage-dependent cell death (31),(32). In contrast, some immortalized cell types such as hMSC-TERT cells may exhibit anchorage-independent growth, as determined by the soft agar colony assay (33). We investigated whether primary hMSC were capable of spreading on the outer membrane of the microcapsule, even when cells were cultured under serum-free conditions (to avoid the presence of serum-derived adhesion molecules such as fibronectin (34)). Interestingly, we found that primary hMSCs were able to spread on the outer microcapsule membrane suggesting that: a) PEC that forms the microcapsule membrane is not cytotoxic, and b) the microcapsule membrane, although not made of natural extracellular matrix components, may provide adhesion signals to primary hMSCs. Interestingly, it has been previously shown that the outer and the inner membrane of the SCS-pDADMAC microcapsules
have similar surface composition (35),(36). Although we found that hMSC-TERT cells were able to spread to some extent on the inner membrane of the microcapsule, we observed that primary hMSCs failed to do so. We hypothesized that anoikis, or anchorage-dependent cell death related to the structure of the microcapsule membrane may not be the main (or the only) cause of the reduced viability in primary cells. Nevertheless, primary hMSCs may require other attachment and spreading factors for long-term survival that the microcapsule membrane may not provide. Immortalized cells may overcome this possibly due to up-regulation of other cell survival mechanism. In this study we have not investigated whether the inner membrane provided attachment signals to the cells (e.g. via integrin expression by encapsulated cells).

Furthermore, membrane pores below a critical size may cause inefficient diffusion and delay in the transport of substances into the microcapsule, which could negatively affect the growth and survival of the encapsulated cells. Membranes formed by SCS and pDADMAC crosslinking are known to have a very low molecular weight cut-off (MWCO). For instance, Zhang et al. showed that SCS-pDADMAC microcapsules prepared with their conditions had a MWCO of ~ 14k Da, meaning that proteins larger than 14 kDa hardly permeate through the membrane (37) Dautzenber et al. characterized the membrane cut-off limits of SCS-pDAMAC microcapsules prepared with different pDADMAC solutions with mass portions of size fractions >10kDa, and they found that the cut-off range corresponded to a molecule with R_h ~ 1.9 nm (equal to a globular protein of about 18k Da) (38). According to Dautzenber et al., size exclusion properties may be influenced by several variables during the capsule formation including the concentration of pDADMAC and its molecular weight, the polymerization time and the presence of monovalent salt in the pDADMAC solution (25),(26),(38). Therefore, in this study, we have characterized the properties of the PEC membrane made with the conditions described in the experimental section. SEM
analysis of the microcapsule membrane showed that this was formed by a very compact structure with no visible large pores. However, the dehydration step performed during the sample preparation could have had a detrimental effect on the structure of the membrane’s pores. Therefore, we studied the microcapsule membrane permeability properties in a functional manner (e.g. by the ability of molecules of different molecular weight to diffuse into the microcapsule). Results from this permeability study showed that only small molecules (i.e. 10 kDa dextran) were able to freely permeate the microcapsule membrane, while 20 kDa dextran permeated only to an extent and BSA (66.5 kDa) appeared absorbed to the outer membrane. Moreover, we demonstrated that large molecules such as antibodies (i.e. 70k Da dextran and 180 kDa IgY) were not able to permeate and therefore, cells could be protected from an antibody-mediated immune attack.

Table 1 shows the correspondent hydrodynamic radius ($R_h$) of some molecules as a reference for permeability estimation. The $R_h$ is employed for comparing the size of proteins, which have different molecular shapes (i.e. dextrans or globular proteins). It is defined as the radius of a hypothetical hard sphere that diffuses with the same speed as the molecule under examination and it is closely related to solvent motility. From the permeability experiments, we can stipulate that the pore size of the microcapsule membrane may correspond, approximately, with the size of a molecule with $R_h$ between 3.2 nm and 3.48 nm. Therefore, we would expect that small molecules such as glucose, most cytokines and growth factors (i.e. VEGF which has a molecular weight of 45 kDa and a calculated $R_h$ of 3.02 nm $^{(39)}$) present in the culture medium and/or released by the encapsulated cells would be able to diffuse through the microcapsule membrane. In this regard, we could observe a release of angiogenic factors, including VEGF, from encapsulated cells after 7 days in culture, as determined by the matrigel tubule formation assay
and quantification of VEGF levels from the conditioned medium. We found that hMSC-TERT cells secreted higher levels of VEGF than primary hMSCs probably due to a higher viability of these encapsulated cells at that chosen time point.

FBS is used as an almost universal growth supplement in cell culture media for most types of human and animal cells. It is composed of low and high molecular weight biomolecules including hormones, vitamins, carbohydrates and proteins including serum proteins, growth factors and cytokines, transport proteins, attachment and spreading factors and enzymes, which are vital for the attachment, growth and proliferation of cells \((34),(40)\). We investigated whether primary hMSCs and hMSC-TERT cells were able to grow in FBS-free medium or in medium depleted of macromolecules \(>50\) kDa (50kD-depleted medium). We found that primary hMSCs were more sensitive to serum-free conditions than MSC-TERT cells, and that they need macromolecules (i.e. \(<50\) kDa) present in the serum for survival and proliferation. For instance, primary hMSCs may need macromolecules such as transferrin (80k Da) (important for transferring iron to the cells \((34)\)) and serum albumin (66.5 kDa) (a potent antioxidant especially in serum-starved conditions \((41),(42)\)) for optimal proliferation and survival. In contrast, the growth and viability of immortalized cell lines such as hMSC-TERT cells was less compromised by these conditions, as these cells proliferated even in the absence of serum.

Based on our results, we concluded that liquid-core microcapsules must have a membrane pore size that allows the entrance of proteins \(>50\) kDa in order to support primary hMSCs viability. However, if immunoprotection of enclosed cells is of interest, these microcapsules must have a pore size \(<150\) kDa (e.g. IgG). Nonetheless, finding a correct balance could be challenging.

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FIGURE LEGENDS

**Figure 1. Chemical and morphological characterization of microcapsules.** A) Schematic representation of microcapsule made of SCS and pDADMAC to encapsulate cells. B) Ultrastructure of microcapsules visualized using Back Scattered Electron Mode (BES) scanning electron microscopy (SEM). C) Microcapsule visualized using standard SEM protocol showing the loss of the liquid core after dehydration process. D) FTIR spectrum of naïve SCS and pDADMAC, and PEC formed after polymerization. E) Representative graph of microcapsule size distribution.

**Figure 2. Behavior of primary hMSCs encapsulated in SCS-pDADMAC microcapsules.** A) CLSM images showing viability of encapsulated cells stained with Calcein AM (green fluorescence indicates living cells) and Ethidium Homodimer-1 (EthD-1, red fluorescence indicates dead cells). Control dead cells were visualized using excitation and emission wavelengths of ~528nm/~617nm (dead cells) and ~495 nm/~515 nm (live cells). B) Metabolic activity using AlamarBlue® assay at days 0, 3, 7 and 14 post-encapsulation. Absorbance was measured at 550 and 595nm. Statistical analysis by one-way ANOVA with Tukey’s multiple comparison test (*p<0.05 and **p<0.01 compared to day 0). C) Orthographic projection of microcapsules showing the localization of hMSCs inside of microcapsules. Cells were stained with Calcein AM and EthD-1. D) Cytoskeleton organization of encapsulated hMSCs stained with Phalloidin-FITC (green) for F-actin and propidium iodide (red) for nucleus.

**Figure 3. Biological response of primary hMSCs to microcapsule material measured by AlamarBlue® assay.** A) Cytotoxicity analysis of microcapsule components to hMSCs by direct contact with sterile empty microcapsules (direct contact), by contact with extracts of the device through diffusion (indirect contact) and by direct contact with 2% and 4% (w/v) SCS at 1:3 and 2:3 ratios (SCS:complete medium). Statistical analysis performed by unpaired Student’s t-test (n.s.). B) CLSM images using transmitted laser light (TL) and LIVE/DEAD dyes (ZP: z projection and OP: orthographic projection) showing spreading of primary hMSC on the outer microcapsule membrane immediately and 24h after cell seeding. Statistical analysis performed by one-way ANOVA with Dunnett’s multiple comparison test (n.s.).

**Figure 4. Behavior of immortalized hMSC-TERT cells encapsulated in SCS-pDADMAC microcapsules.** A) CLSM images showing viability of encapsulated cells stained with Calcein
AM (green fluorescence indicates live cells) and Ethidium Homodimer-1 (EthD-1, red fluorescence indicates dead cells). B) Metabolic activity using AlamarBlue® assay at days 0, 3, 7 and 14 post-encapsulation. Absorbance was measured at 550 and 595nm. Statistical analysis by one-way ANOVA with Tukey’s multiple comparison test (n.s.). C) Orthographic projection of microcapsules showing the localization of hMSC-TERT cells inside of microcapsules. Cells were stained with Calcein AM and EthD-1. D) Cytoskeleton organization of encapsulated hMSC-TERT cells stained with Phalloidin-FITC (green) for F-actin and propidium iodide (red) for nuclei.

**Figure 5. Diffusion properties of microcapsule membrane.** A) SEM images of microcapsule membrane. Left image is the surface of microcapsule membrane; center image is a higher magnification of left image; right image shows a transverse section of a microcapsule membrane. B) CLSM images of microcapsules incubated with FITC- labeled dextrans of 10, 20 and 70kDa (with hydrodynamic radius $R_h$ of 2.3 nm, 3.2 nm and 5.8 nm, respectively), FITC-coupled BSA (66.5 kDa, $R_h$ 3.48 nm) and FITC-coupled chicken immunoglobulin IgY (180 kDa, $R_h$>5.5 nm).

**Figure 6. Influence of macromolecular components from culture medium on primary hMSCs and hMSC-TERT growth.** A) Protein composition of serum-free medium (lane 1), complete medium (lane 2), 50K-depleted medium (lane 3) and 50K-concentrated medium (lane 4) by SDS-PAGE and silver staining. B) Growth curve of primary hMSCs and hMSC-TERT cells cultured in complete medium —■—, serum-free medium --○--, 50K-depleted medium ⋯▲⋯ and 50K-concentrated medium ---- ▼---. C) Fold-change of primary hMSCs (**$p<0.01$ significance compared to complete medium) and hMSC-TERT growth (***$p<0.001$ compared to complete medium; +++$p<0.001$ compared to serum-free medium and 50K-depleted medium) at day 9 compared to day 0. Dotted line at 1 represents baseline (day 0). Statistical significance by one-way ANOVA with Tukey’s multiple comparison test.

**Figure 7. In vitro angiogenic potential of encapsulated cells.** A) Images of HUVECs tubule formation in matrigel after 14h incubation with EGM (positive control) EBM (negative control) and conditioned media from encapsulated hMSC-TERT cells and primary hMSCs 7 days post-encapsulation. B) Quantification of tubule-like structures 7 days post-encapsulation. Results are represented as fold-change compared to EGM. Statistical analysis by one-way ANOVA with Tukey’s multiple comparison test (*$p<0.05$ and **$p<0.01$ significance compared to EBM,
+++p<0.001 significance compared to EGM). C) Quantification of VEGF levels released from encapsulated cells 7 days post-encapsulation over a period of 24h, 48h and 72h by ELISA (unpaired Student’s t-test *p<0.05 and **p<0.01 significance of hMSC-TERT compared to primary hMSC; two-way ANOVA with Tukey’s test +p<0.05 significance of hMSC-TERT at 72h compared to 24h).

**Supporting information.** Three-dimensional simulation of microcapsule containing primary hMSCs and hMSC-TERT cells (Video S1 and Video S2), characterization of primary hMSC by flow cytometry and tri-lineage differentiation (Figure S1) morphology of primary hMSCs and hMSC-TERT cells under CLSM (Figure S2), viability of other cell types (Figure S3), fluorescence intensity spectrum (Figure S4) and quantification of VEGF levels from non-encapsulated cells (Figure S5).
List of Figures:

Figure 1
Figure 2
Figure 3

A

i) Cytotoxicity by Direct Contact

![Graph showing cytotoxicity by direct contact with hMSCs and hMSCs + Caps]

ii) Cytotoxicity by Indirect Contact

![Graph showing cytotoxicity by indirect contact with hMSCs and hMSCs + CM]

iii) Cytotoxicity of SCS by Direct Contact

![Graph showing cytotoxicity of SCS with hMSCs and ratios 1:3, 2:3, 1:3, 2:3 with 2% (w/v) and 4% (w/v) SCS]

B

![Images of TL, ZP, and OP immediately and 24h after treatment]
Figure 4

(A) Images showing cellular morphology at various time points. (B) Bar graph depicting metabolic activity over different days post-encapsulation. (C) Additional images illustrating cellular structure. (D) Further visual data supporting the findings.
Figure 5

A

Dextran-10kDa (2.3nm)
Dextran-20kDa (3.2nm)
BSA- 66.5kDa (3.48nm)
IgY-180kDa (>5.5nm)
Dextran-70kDa (5.8nm)

Capsule membrane

B

Good for immunoprotection
Figure 6

A

B

Primary hMSCs

hMSC-TERT

C

Primary hMSCs (Day 9)

hMSC-TERT (Day 9)
Figure 7

A

B

C

Fold change compared to EGM

Primary hMSCs

Human MSC-TERT

Incubation time (h)

VEGF (pg/ml)