Isolation and Characteristics of Mesenchymal Stromal Cells from Different Parts of Placenta

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Abstract

Placenta performs the following functions: protection, nutrition, respiration, hormone production and excretion. As it is a great source of different cells, we are more and more interested to isolate them from the placenta. Cells can be harvested by non-invasive methods and without any ethical concerns. Due to its structure placenta contains Mesenchymal Stromal Cells (MSCs) of maternal and fetal origin. To make the selection easy, in our experiment we only used placentas from women who gave birth to boys. We used two methods to isolate MSCs (CD - collagenase digestion and MC - mechanical cut) from different parts of the placenta: amnion, chorion, villi and decidua basalis.

MSCs were of CD73⁺, CD90⁺, CD105⁺, CD 14⁻, CD19⁻, CD45⁻, HLA-DR⁻ cell surface phenotype, adherent and capable to differentiate into osteocytes, adipocytes, chondrocytes. These data fulfilled minimal characterization criteria of MSCs.

We can isolate fetal and maternal MSCs from placenta. The origin of isolated cells was tested with the use of Fluorescence in situ Hybridization (FISH). MSCs isolated from the same placenta from one tissue can show different origins, for example, MSCs from chorion isolated by MC show maternal origin, but MSCs from the same amnion isolated using another method (CD) show fetal origin.

A placenta mostly consists of fetal-derived cells. However, close contact between fetal (amnion, chorion and villi) and maternal (decidua basalis) parts is responsible for presence of the maternal cells in the fetal part and fetal cells in the maternal part of a placenta. Isolation of pure: maternal or fetal MSCs from the placental tissue allows better characterization of MSC-based product for clinical purposes. If we are able to produce a pure population of maternal MSCs, we will gain the ability to apply a more personalized therapy for the mother.

Keywords: Mesenchymal stromal cells; MSC; Placenta; Cell therapy

Introduction

Today, there is a new field in the modern biomedicine the cell therapy, in which in vivo stem cells are transplanted to compensate for tissue dysfunction and regenerate damaged organs. Stem cells have two main features: self-renewal and the ability to differentiate into other cell types. They are a pool of undifferentiated progenitor cells of various types. The use of stem cells is the most promising direction of the cell therapy.

Stem cells are classified into embryonic and adult stem cells. Embryonic Stem Cells (ESCs) have a high potential for differentiation into many types of cells. The use of ESCs raises ethical questions and is associated with a high risk of cancer development. Additionally, ESCs express HLA, which leads to transplant rejection. In contrast, adult stem cells, for example mesenchymal stromal cells and hematopoietic stem cells can be isolated without any ethical problems using a non-invasive method.

Mesenchymal Stromal Cells (MSCs) are a good source for the cell therapy thanks to their properties. It was demonstrated that MSCs have the ability of self-renewal, secret factors that can facilitate tissue repair, and can differentiate into different types of cells, such as chondrocytes, adipocytes, osteocytes, cardiomyocytes, neuronal cells and other [1]. MSCs exert very important immunomodulatory effects: they suppress T- and B-cell proliferation and natural killer cells function, and they also limit the expression of the Major Histocompatibility Complex II (MHC II) [2,3]. Thanks to these properties, MSCs can be used in an effective therapy of the Graft-Versus-Host Disease (GVHD) [4,5]. It was shown that MSCs migrate to the sites of tissue injury (Table 1). “Young” MSCs isolated from the placenta show a better proliferation and differentiation ability than “adult” MSCs [6]. For this reason, they have been used in a number of clinical trials (www.clinicaltrials.gov).

Friedenstein was the first to isolate and describe mesenchymal stromal cells from bone marrow [7]. It is known that the number of MSCs in the body, as well as their ability to proliferate and differentiate decline significantly with age [8].

The placenta is a very good source of a range of cells and hence has been attracting a growing interest. Cells can be harvested by non-invasive methods and without any ethical problems. Due to its structure, the placenta contains MSCs of maternal and fetal origin (Figure 1). The decidua basalis is a part of the endometrium adjacent to the myometrium. The decidua basalis is the best supplied with maternal blood and later expands to form the maternal part of the placenta. MSCs are also found in the amnion and the chorion - two fetal membranes,
and in the chorionic villi. We attempted to isolate MSCs of maternal and fetal origin from each of these parts of the placenta. If we are able to produce a pure population of maternal MSCs, we will gain the ability to apply a more personalized therapy for the mother.

**Methods**

**Term placenta sample collection**

Human full-term placentas were obtained from healthy women aged 19-33 years, at the time of a routine caesarean section or vaginal delivery in an affiliated hospital in Cracow. The mean gestational age was 39 weeks (37-41 weeks). To facilitate the selection of donors, in our experiment we only used placentas of women who gave birth to boys. Placentas were collected in accordance with a protocol approved by the Committee of the Ministry of Health in Poland. Placentas were only used from 1 or 2 passages.

**Cell isolation**

After dissection, tissue samples were washed with sterile PBS (phosphate buffered saline, pH 7.4, ice-cold) and treated with antibiotics. Cells were isolated using two methods: Mechanical Cut (MC) and collagenase digestion (CD). Samples were cut into pieces, washed with PBS twice and incubated at 37°C in 75 cm² flasks in Mesencult (STEMCELL Technologies) with supplement and antibiotic/antimicotic (Gibco) in 25 cm² flasks (BD Falcon), and cultured at 37°C in 5% CO₂ and 90% of humidity. In the CD method, the cells were harvested after one to seven days of culturing. We used cells only from 1 or 2 passages.

**Flow cytometry**

For phenotypic evaluation, all the MSCs extracted from culture were incubated for 30 min. with phycoerythrin-conjugated antibodies against the human antigens: CD73 (BD Biosciences), CD90 (BD Pharmingen, BD Biosciences), CD105 (BD Pharmingen, BD Biosciences), and fluoroscin isothiocyanate-conjugated antibodies against the human antigens: HLA-DR, CD34, CD45, CD19, CD14 (BD Bioscience). Samples were analyzed in a FACS Calibur machine (BD Biosciences). The minimal criteria for MSCs were: adherence to plastic; positive expression of CD73, CD90 and CD105; lack of expression of CD45, CD34, CD14, CD19 and HLA-DR; and the ability to differentiate into osteocytes, chondrocytes and adipocytes.

**Fluorescence in situ hybridization (FISH)**

Thirty seven fresh placental tissue samples (chorion, amnion, red tissue of placenta and separated villi fiber), after manual preparation and in vitro culture, were harvested according to standard cytogenetics procedures. After cell synchronization by colcemid for 20 min at 37°C (10 µg/ml, Biosera), pellet cells undergo a hypotonic treatment using 0.075 M KCL solution (Merck) for 15 minutes at 37°C to swell the cells. The cells were then fixed in cold Carnoy’s fixative solution composed 3:1 methanol and 100% acetic acid (Merck) washed three times to ensure complete removal of cytoplasmic debris. The resulting suspension of metaphase and interphase cells was applied to microscopic slides. FISH was performed with the commercially available probe SE X(DXZ1)/Y (DYZ3) (Kreatech Diagnostics), dedicated for identification of aneuploidy. The procedure was applied according to the manufacturer’s protocol. Slides were analyzed using an epifluorescence microscope Imager. Z2 (Carl Zeiss) and documented using an ISIS (Metasystems) Imaging System.

**Results**

We successfully were able to isolate MSCs from different parts of placenta. Morphologically, MSCs were fibroblast-like cells. We have not rotation at 37°C with 1% collagenase IV (Gibco, Life Technologies) for 1.5 hours (CD). After collagenase digestion, cells were collected by centrifugation at 300 rpm for 7 min., followed by washing with DMEM with 10% FBS two times. Then the CD cells were resuspended in Mesencult (STEMCELL Technologies) with supplement and antibiotic/antimicotic (Gibco) in 25 cm² flasks (BD Falcon), and cultured at 37°C in an atmosphere with 5% CO₂ and 90% of humidity. In the MC method, after one month of culturing the cells migrated from the tissues and adhere the flask surface. In the CD method, the cells were harvested after one to seven days of culturing (Figure 2). We used cells only from 1 or 2 passages.

**Table 1:** Selected properties of MSCs which can be useful in the stem cell therapy.

<table>
<thead>
<tr>
<th>Function</th>
<th>Literature</th>
</tr>
</thead>
<tbody>
<tr>
<td>INJURY</td>
<td>Tropism for sites of tissue injury</td>
</tr>
<tr>
<td>INFLAMMATION</td>
<td>T-lymphocyte activation, macrophage infiltration</td>
</tr>
<tr>
<td>NEUROGENESIS</td>
<td>Neuronal growth and differentiation</td>
</tr>
<tr>
<td>SECRETION of factors</td>
<td>Secretion of angiogenic and neurotrophic factors</td>
</tr>
<tr>
<td>APOPTOSIS</td>
<td>Apoptosis (apoptotic cell death)</td>
</tr>
<tr>
<td>NEURAL SYSTEM</td>
<td>Proliferation and activation of astrocytes</td>
</tr>
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</table>

and in the chorionic villi. We attempted to isolate MSCs of maternal and fetal origin from each of these parts of the placenta. If we are able to produce a pure population of maternal MSCs, we will gain the ability to apply a more personalized therapy for the mother.

**Methods**

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noticed a significant morphological difference between cells isolated from different parts and by different methods. Process of differentiation of investigated cells into chondrocytes, adipocytes and osteocytes also looks very similar despite the part of placenta they came from.

**Flow cytometry**

Using cytometry, we searched for the following surface antigens of MSCs: CD73⁺, CD90⁺, CD105⁺, CD14⁻, CD19⁻, CD34⁻, CD45⁻ and HLA-DR⁻. Every part of placenta shows the same immunophenotype (Figure 3).

**Differentiation**

The extracted MSCs were able to differentiate into osteocytes, adipocytes and chondrocytes (Figure 4). The differentiation of MSCs from different parts of placenta follows the same pattern. Therefore, they fulfilled the minimal characterization criteria of MSCs. Exemplary differentiations of the cells from *decidua basalis* are presented in Figure 4.

**Fluorescence in situ analysis (FISH)**

The origin of the isolated cells was tested using FISH method. Fluorescence *in situ* hybridization was performed mostly on interphase cells, and additionally on metaphase cells, to detect the number of copies of highly repetitive satellite DNA sequences located in the pericentric heterochromatin of chromosomes X and Y. Between 107 to 728 cells were analysed per every sample. The FISH tests results are presented in Table 2.

In this experiment, we demonstrated that MSCs isolated from the same placenta and the same tissue can be of different origin, for example MSCs isolated mechanically (MC) from the chorion were of maternal origin, while MSCs from the same chorion sample isolated with the use of the Collagenase Method (CD) had fetal origin. MSCs isolated mechanically from the fetal part (chorion) of the placenta showed maternal contamination. MSCs from the amnion isolated by both methods showed maternal origin/contamination. It should be noted that the maternal part of the placenta (*decidua basalis*) had fetal contamination in the cells isolated mechanically (Figure 5). MSCs isolated by both CD and MC had different origin (Figure 6) and in the some cases – contamination by maternal or fetal cells.

The graphs show MSCs of maternal, fetal or mixed (maternal-fetal or fetal-maternal) origin by isolation group (CD or MC). MSCs from the chorion isolated by CD had more cells of maternal origin than MSCs isolated mechanically, while MSCs from the same tissue were generally of fetal origin. However, MSCs from chorionic villi and amnion isolated both by CD and MC had mostly fetal origin.

**Discussion and Conclusion**

The placenta consists of a larger fetal part (the membranes: chorion and amnion, villi) and a maternal part – *decidua basalis*. The placenta performs the following functions for the fetus: protection, nutrition, respiration, hormone production and excretion. The placenta grows throughout all pregnancy.

In the present study, we isolated fetal and maternal population of stem cells from different parts of the placenta. All the isolated MSCs had typical fibroblastic morphology, expressed cell-surface markers (CD73, CD90, CD105) and differentiated into a mesodermal lineage, a criterion recommended by the International Society of Cell Therapy.

Several protocols described two different methods of isolation: mechanical separation and enzyme digestion [9,10]. In our experiment, we isolated MSCs by mechanical cut and collagenase digestion to compare cell populations from a given tissue isolated with different methods. The enzymatic method, MSCs can be obtained faster.

We successfully differentiated isolated cells into mesodermal lineage cells: osteocytes, chondrocytes and adipocytes.

A placenta mostly consists of fetal-derived cells. However, a close contact between fetal (amnion, chorion and villi) and maternal (*decidua basalis*) parts will lead to the presence of maternal cells in the fetal part.
and fetal cells in the maternal part of the placenta. Our controversial FISH-results require further study and analysis.

During pregnancy maternal cells migrate into the fetus and fetal cells migrate into the mother [11,12]. Presence of maternal cells within the fetus: in the liver, skin, thymus [13]. Suggested that placenta is not a good barrier than previously thought. Maternal cells must to have ability to proliferate, because they have been found in adults [12]. In 2008 Chen and his team suggested that placental cells migration is connected with vascular endothelial growth factor A (VEGF A) [14]. Galazios et al. demonstrated that VEGF A concentration higher in the fetal circulation than in maternal that probably promote cell trafficking through placenta [15]. Nijagal et al. shown that inflammatory disease (autoimmune processes, complication during pregnancy, congenital anomalies and others) [16] during pregnancy changes cell trafficking [17]. Similar to maternal cells, fetal cells were detected into different maternal organs and tissues: liver, kidney, bone marrow [18]. In our study all mothers have cytomegalovirus (CMV), and some authors proofed that CMV can promote immune cell migration and increase the level of VEGF A [19,20], which probably promote placental cell migration.

Figure 3: Exemplary flow cytometric histogram plots after immunopheno typing of placenta-derived MSCs from amnion. Transparent histograms represents cells population stain with polyclonal isotype control antibodies conjugated to fluorescein (FITC) or phycoerythrin (PE) (green or red, respectively). Filled histograms represent cell population labelled with monoclonal antibodies against CD34 (A), CD45 (B), HLA-DR (C), CD14 (D), CD19 (E), CD73 (F), CD90 (G), and CD105 (H) conjugated with fluorescein or phycoerythrin (green or red, respectively).
less labor-consuming. Despite previous publication, decidua basalis is not the only part of the placenta which is rich in maternal cells. Villi, as well as amnion and chorion also contain maternal cells. Isolation of pure: maternal or fetal MSCs from the placental tissue allows better characterisation of MSC-based product for clinical purposes. Now there is no evidence that the pure population of maternal or fetal mesenchymal stem cells has better therapeutic capabilities. Probably, the banking and followed by clinical application of mixed population of maternal and fetal MSCs will potentially lead to a greater and interesting therapeutic effect. Further research is needed to verify this hypothesis [23-31].

Acknowledgments

Some scientists suggested that fetal MSCs populations are contaminated by maternal cells that prevail after successive passages [21]. Others indicated that a population of fetal cells can be isolated without maternal contamination [9,10,22]. The possibility of isolation of a pure population of maternal or fetal MSCs depends on a wide variety of factors, such as the mother’s diseases, etc.

Our research confirms that the placenta is an unique material, which contains MSCs of different origin: maternal and fetal. Both: collagenase digestion and mechanical cutting resulted in high amount of cells harvested, but usage of enzyme makes this process faster and
The authors acknowledge the help of Kurilovich Ludmila for placenta’s picture.

References


